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(54) Title: CRYSTALLOGRAPHIC STRUCTURE OF THE ANDROGEN RECEPTOR LIGAND BINDING DOMAIN

(57) Abstract: The first crystal structure of the androgen receptor ligand binding domain has been determined to 2.0 angstrom resolution. Disclosed are the coordinates for the crystal structure, and methods for determining agonists, partial agonists, antagonists, partial antagonists, and selective androgen receptors modulators (SARMS) of the androgen receptor.

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CRYSTALLOGRAPHIC STRUCTURE OF THE ANDROGEN RECEPTOR LIGAND BINDING DOMAIN

Field of Invention

The present invention relates to compositions and crystals of androgen receptor ligand binding domain optionally in complex with its ligand. This invention also relates to methods of using the structure coordinates of the androgen receptor ligand binding domain /ligand complex to solve the structure of similar or homologous proteins or protein complexes. This invention also relates to methods for designing and selecting ligands that bind to the androgen receptor and methods of using such ligands.

Background of the Invention

The androgen receptor (AR) is a member of the steroid nuclear-receptor superfamily of ligand-dependent transcription factors. The binding of androgen to AR initiates the gene activation required for male sex development.

AR is an important target primarily in two drug discovery areas. In oncology drug discovery, inhibitors (antagonists or partial antagonists) of androgen receptor function are useful for treatment of anti-androgen refractory prostate cancer. In metabolic diseases drug discovery, agonists or partial agonists to the androgen receptor in muscle are useful to treat age-related diseases.

As with the other members of the steroid receptor family, AR has several functional domains including a DNA binding domain (DBD), and a 261 residue ligand-binding domain (LBD) (Mw = 30,245 Da) which contains the androgen binding site, and is responsible for switching on the androgen function.

Development of synthetic ligands that specifically bind to androgen receptors has been largely guided by trial and error method of drug design despite the importance of the androgen receptor in physiological processes and medical conditions such as prostate cancer and modulation of reproductive organ modulation. Previously, new ligands specific for androgen receptors were discovered in the absence of information on the three dimensional structure of the androgen receptor with a bound ligand. Before the present invention, researchers were

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essentially discovering androgen receptor ligands by probing in the dark and without the ability to visualize how the amino acids of the androgen receptor held a ligand in its grasp.

Consequently, it would be advantageous to devise methods and compositions for reducing the time required to discover ligands to the androgen receptor, synthesize such compounds and administer such compounds to organisms to modulate physiological processes regulated by the androgen receptor.

The cDNA and amino acid sequences of human and rat androgen receptors have been described (Proc. Natl. Acad. Sci. U.S.A. 1988 85: 7211-7215). However, there have been no crystals reported of any androgen receptor. Thus, x-ray crystallographic analysis of such proteins has not been possible.

We have discovered the first crystal structure of the androgen receptor ligand binding domain (AR-LBD). Our understanding or the androgen receptor structure has allowed for the determination of the ligand binding site for selective androgen receptor modulators (SARMs).

Summary of the Invention

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The present invention provides crystals of AR-LBD and crystals of an AR-LBD bound to a ligand, i.e. an AR-LBD/AR-LBD ligand complex. Most preferably the AR-LBD ligand is dihydrotestosterone (DHT). Thus, the present invention is directed to a crystal of an AR-LBD comprising:

- 1) an AR-LBD and an AR-LBD ligand or
- 2) an AR-LBD without an AR-LBD ligand; wherein said crystal diffracts to at least 3 angstrom resolution and has a crystal stability within 5% of its unit cell dimensions. The crystal of AR or AR-LBD preferably has at least 200 amino acid and preferably comprises amino acid sequence 672 to 917 of rat AR or the AR amino acid sequence 672 to 917 of human AR.

The present invention also provides the structure coordinates of the AR-LBD/AR-LBD ligand complex. The complete coordinates are listed in Table A.

The present invention also provides a method for determining at least a portion of the three-dimensional structure of molecules or 35 molecular complexes which contain at least some structurally similar

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features to the androgen receptor ligand binding domain. It is preferred that these molecules or molecular complexes comprise at least a part of the ligand binding site defined by structure coordinates of AR-LBD amino acids V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877 and F878 according to Table A, or a mutant or homologue thereof. Since the protein sequences for rat and human AR LBD are identical, the human numbering system was used herein.

The present invention also provides a machine-readable data storage medium which comprises a data storage material encoded with machine readable data defined by the structure coordinates of an AR-LBD/AR-LBD ligand or ligand complex according to Table A or a homologue of the complex.

The present invention further provides a binding site in AR-LBD for an AR-LBD ligand as well as methods for designing or selecting AR modulators including agonists, partial agonists, antagonists, partial antagonists and/or selective androgen receptor modulators (SARMs) of AR using information about the crystal structures disclosed herein.

Brief Description of the Drawing

Figure 1 is a ribbon style drawing of the Androgen Receptor LBD. The substrate DHT is shown as a ball-and-stick figure.

Figure 2 is a comparison of the androgen receptor ligand binding domain with progesterone receptor ligand binding domain.

Figure 3 provides three views of the omit electron density map of dihydrotestosterone (DHT) in the hormone-binding site of AR-LBD. There are hydrogen bonds between the steroid and the side chains of Arg 752 and Asn 705.

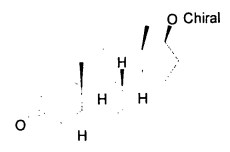
Figure 4 is a comparison of the binding of dihydrotestosterone to AR-LBD (top) and of progesterone to PR-LBD (bottom). Note that an additional hydrogen bond interaction would be possible if both the sidechains of both N719 and the progesterone were flipped.

Detailed Description of the Invention

The first crystal structure of the androgen receptor ligand binding domain (AR-LBD) has been determined to 2.0 Å resolution.

Crystals of rat AR-LBD were grown from precipitating solutions containing 0.9 M Sodium Tartrate, 0.1 M Na Hepes, pH 7.5. X-ray diffraction from the crystals have the symmetry and systematic absences of the orthorhombic space group P212121 with unit cell dimensions $a = 56.03 \, \text{Å}$, $b = 66.27 \, \text{Å}$, $c = 70.38 \, \text{Å}$, and one molecule per asymmetric unit (Mathews Volume = $2.16 \, \text{Å}^3 \, \text{Da}^{-1}$). The structure was determined by the method of molecular replacement using the structure of the Progesterone Receptor LBD (PR-LBD) as the search model.

The complex of AR-LBD with dihydrotestosterone (DHT) shows
the mode of binding of the steroid to the receptor in the agonist
conformation.



Dihydrotestosterone

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The following abbreviations are used throughout the application:

A = Ala = Alanine

V = Val = Valine

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I = Ile = Isoleucine

P = Pro = proline

F = Phe = phenylalanine

W = Trp = Tryptophan

25 M = Met = Methionine

G = Gly = Glycine

S = Ser = Serine

T = Thr = Threonine

C = Cys = Cysteine

30 Y = Tyr = Tyrosine

N =Asn = Asparagine

O =Gln = Glutamine

D = Asp = Aspartic Acid

E = Glu = Glutamic Acid

K = Lys = Lysine

R = Arg = Arginine

H = His = Histidine

"Atom type" refers to the element whose coordinates have been determined. Elements are defined by the first letter in the column.

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"X, Y, Z" crystallographically define the atomic position determined for each atom.

"B" is a thermal factor that measures movement of the atom around its atomic center. 15

"Occ" is an occupancy factor that refers to the fraction of the molecules in which each atom occupies the position specified by the coordinates. A value of "1" indicates that each atom has the same conformation, i.e., the same position, in all molecules of the crystal.

Additional definitions are set forth in the specification where necessary.

The androgen receptor (AR) described herein is intended to include any polypeptide which has the activity of the naturally occurring androgen receptor. The AR and AR-LBD contemplated herein includes all vertebrate and mammalian forms such as rat, mouse, pig, goat, horse, guinea pig, rabbit, monkey, orangutan and human. Such terms also include polypeptides that differ from naturally occurring forms of AR and AR-LBD by having amino acid deletions, substitutions, and additions, but which retain the activity of AR and AR-LBD, respectively. The crystal structure of the invention preferably contains at least 25%, more preferably at least 50%, more preferably at least 75%, more preferably at least 90%, more preferably at least 95%, more preferably at least 99%, and most preferably all of the coordinates listed in Table A. The crystal of the AR-LBD/AR-LBD ligand of the invention preferably

has the following unit cell dimensions in angstroms: $a = 56.03 \pm 5\%$, b

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= $66.27 \pm 5\%$, $c = 70.38 \pm 5\%$ and an orthorhombic space group P212121.

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The AR-LBD ligand of this invention is any peptide, peptide mimetic or nonpeptide, including small organic molecules, that is capable of acting as a ligand for AR-LBD. In a preferred embodiment, the AR-LBD ligand is an AR modulator. By "AR modulator" it is meant an agonist or activator, a partial agonist or partial activator, an antagonist or inhibitor, or a partial antagonist or partial inhibitor which demonstrates tissue specific activations of the AR. Such compounds are also referred to herein as SARMs (selective androgen receptor modulators) of the AR-LBD. Examples of preferred agonists include androgens such as dihydrotestosterone.

The peptides referred to herein (e.g., AR, AR-LBD, and the like) may be produced by any well-known method, including synthetic methods, such as solid phase, liquid phase and combination solid phase/liquid phase syntheses; recombinant DNA methods, including cDNA cloning, optionally combined with site directed mutagenesis; and/or purification of the natural products, optionally combined with enzymatic cleavage methods to produce fragments of naturally occurring

Advantageously, the crystallizable compositions provided by this invention are amenable to x-ray crystallography. Thus, this invention also provides the three-dimensional structure of the AR-LBD/AR-LBD ligand complex, particularly the complex of rat AR-LBD with dihydrotestosterone.

The three-dimensional structure of the AR-LBD / dihydrotestosterone complex of this invention is defined by a set of structure coordinates as set forth in Table A. The term "structure coordinates" refers to Cartesian coordinates derived from mathematical equations related to the patterns obtained on diffraction of a monochromatic beam of X-rays by the atoms (scattering centers) of an androgen receptor/dihydrotestosterone complex in crystal form. The diffraction data are used to calculate an electron density map of the repeating unit of the crystal. The electron density maps are then used to establish the positions of the individual atoms of the complex.

Those of skill in the art will understand that a set of structure coordinates for a receptor or receptor/ligand complex or a portion

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thereof, is a relative set of points that define a shape in three dimensions. Thus, it is possible that an entirely different set of coordinates could define a similar or identical shape. Moreover, slight variations in the individual coordinates will have little effect on overall shape.

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The variations in coordinates discussed above may be generated because of mathematical manipulations of the structure coordinates. For example, the structure coordinates set forth in Table A could be manipulated by crystallographic permutations of the structure coordinates, fractionalization of the structure coordinates; integer additions or subtractions to sets of the structure coordinates, inversion of the structure coordinates or any combination of the above.

Alternatively, modifications in the crystal structure due to mutations, additions, substitutions, and/or deletions of amino acids, or other changes in any of the components that make up the crystal could also account for variations in structure coordinates. If such variations are within an acceptable standard error as compared to the original coordinates, the resulting three-dimensional shape is considered to be the same.

Various computational analyses are therefore necessary to determine whether a molecule or molecular complex or a portion thereof is sufficiently similar to all or parts of the androgen receptor/dihydrotestosterone described above as to be considered the same. Such analyses may be carried out in current software applications, such as the Molecular Similarity application of QUANTA (Molecular Simulations Inc., San Diego, CA) version 4.1, and as described in the accompanying User's Guide.

The Molecular Similarity application permits comparisons between different structures, different conformations of the same structure, and different parts of the same structure. The procedure used in Molecular Similarity to compare structures is divided into four steps:

1) load the structures to be compared; 2) define the atom equivalences in these structures; 3) perform a fitting operation; and 4) analyze the results.

Each structure is identified by a name. One structure is identified as the target (i.e., the fixed structure); all remaining structures

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are working structures (i.e., moving structures). Since atom equivalency within QUANTA is defined by user input, for the purpose of this invention we will define equivalent atoms as protein backbone atoms (N, Cs, C and O) for all conserved residues between the two structures being compared. We will also consider only rigid fitting operations.

When a rigid fitting method is used, the working structure is translated and rotated to obtain an optimum fit with the target structure. The fitting operation uses an algorithm that computes the optimum translation and rotation to be applied to the moving structure, such that the root mean square difference of the fit over the specified pairs of equivalent atom is an absolute minimum. This number, given in angstroms, is reported by QUANTA.

For the purpose of this invention, any molecule or molecular complex that has a root mean square deviation of conserved residue backbone atoms (N, C α , C, O) of less than 1.5 Å when superimposed on the relevant backbone atoms described by structure coordinates listed in Table A are considered identical. More preferably, the root mean square deviation is less than 1.0 Å. In a preferred embodiment of the present invention, the molecule or molecular complex comprises at least a portion of the ligand binding site defined by structure coordinates of AR-LBD amino acids V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877 and F878 according to Table A, or a mutant or homologue of said molecule or molecular complex. More preferred are molecules or molecular complexes comprising all or any part of the ligand binding site defined by structure coordinates of AR-LBD amino acids N705, Q711, R752, F764 and T877 according to Table A, or a mutant or homologue of said molecule or molecular complex. Since the protein sequences for rat and human AR LBD are identical, the human numbering system has been used herein.

The term "complex" or "molecular complex" means AR-LBD or a mutant or homologue of AR-LBD in a covalent or non-covalent association with a chemical entity or compound.

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For purposes of the present invention, by "at least a portion of" it is meant all or any part of the ligand binding site defined by these structure coordinates.

By "mutant or homologue" as used herein it is meant a molecule or molecular complex having a similar structure and/or sequences to AR-LBD. By "similar structure" it is meant a mutant or homologue having a binding pocket that has a root mean square deviation from the backbone atoms of said AR-LBD amino acids of not more than 1.5 Angstroms. By "similar sequence" it is meant a mutant or homologue having 30%, or more preferably 75%, identity with AR-LBD.

The term "root mean square deviation" means the square root of the arithmetic mean of the squares of the deviations from the mean. It is a way to express the deviation or variation from a trend or object. For purposes of this invention, the "root mean square deviation" defines the variation in the backbone of a protein or protein complex from the relevant portion of the backbone of the AR portion of the complex as defined by the structure coordinates described herein.

Once the structure coordinates of a protein crystal have been determined they are useful in solving the structures of other crystals.

Thus, in accordance with the present invention, the structure coordinates of an androgen receptor/dihydrotestosterone complex, and in particular a complex, and portions thereof is stored in a machine-readable storage medium. Such data may be used for a variety of purposes, such as drug discovery and x-ray crystallographic analysis or protein crystal.

Accordingly, in one embodiment of this invention is provided a machine-readable data storage medium comprising a data storage material encoded with the structure coordinates set forth in Table A.

One embodiment utilizes System 10 as disclosed in WO 98/11134, the disclosure of which is incorporated herein by reference in its entirety

For the first time, the present invention permits the use of structure-based or rational drug design techniques to design, select, and synthesize chemical entities, including inhibitory and stimulatory compounds that are capable of binding to AR-LBD, or any portion thereof.

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One particularly useful drug design technique enabled by this invention is iterative drug design. Iterative drug design is a method for optimizing associations between a protein and a compound by determining and evaluating the three-dimensional structures of successive sets of protein/compound complexes.

Those of skill in the art will realize that association of natural ligands or substrates with the binding pockets of their corresponding receptors or enzymes is the basis of many biological mechanisms of action. The term "binding pocket" as used herein, refers to a region of a molecule or molecular complex, that, as a result of its shape, favorably associates with another chemical entity or compound. Similarly, many drugs exert their biological effects through association with the binding pockets of receptors and enzymes. Such associations may occur with all or any parts of the binding pockets. An understanding of such associations will help lead to the design of drugs having more favorable associations with their target receptor or enzyme, and thus, improved biological effects. Therefore, this information is valuable in designing potential ligands or inhibitors of receptors or enzymes, such as inhibitors of AR.

The term "associating with" refers to a condition of proximity between chemical entities or compounds, or portions thereof. The association may be non-covalent -- wherein the juxtaposition is energetically favored by hydrogen bonding or van der Waals or electrostatic interactions -- or it may be covalent.

In iterative drug design, crystals of a series of protein/compound complexes are obtained and then the three-dimensional structures of each complex is solved. Such an approach provides insight into the association between the proteins and compounds of each complex. This is accomplished by selecting compounds with inhibitory activity, obtaining crystals of this new protein/compound complex, solving the three dimensional structure of the complex, and comparing the associations between the new protein/compound complex and previously solved protein/compound complexes. By observing how changes in the compound affected the protein/compound associations, these associations may be optimized.

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In some cases, iterative drug design is carried out by forming successive protein-compound complexes and then crystallizing each new complex. Alternatively, a pre-formed protein crystal is soaked in the presence of an inhibitor, thereby forming a protein/compound complex and obviating the need to crystallize each individual protein/compound complex.

As used herein, the term "soaked" refers to a process in which the crystal is transferred to a solution containing the compound of interest.

The structure coordinates set forth in Table A can also be used to aid in obtaining structural information about another crystallized molecule or molecular complex. This may be achieved by any of a number of well-known techniques, including molecular replacement.

The structure coordinates set forth in Table A can also be used for determining at least a portion of the three-dimensional structure of molecules or molecular complexes which contain at least some structurally similar features to AR. In particular, structural information about another crystallized molecule or molecular complex may be obtained. This may be achieved by any of a number of well-known techniques, including molecular replacement.

Therefore, in another embodiment this invention provides a method of utilizing molecular replacement to obtain structural information about a crystallized molecule or molecular complex whose structure is unknown comprising the steps of:

- a) generating an X-ray diffraction pattern from said crystallized molecule or molecular complex;
 - b) applying at least a portion of the structure coordinates set forth in Table A to the X-ray diffraction pattern to generate a three-dimensional electron density map of the molecule or molecular complex whose structure is unknown; and
 - c) using all or a portion of the structure coordinates set forth in Table A to generate homology models of AR-LBD or any other nuclear hormone receptor ligand binding domain.

Preferably, the crystallized molecule or molecular complex is obtained by soaking a crystal of this invention in a solution.

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By using molecular replacement, all or part of the structure coordinates of the AR-LBD/AR-LBD ligand complex provided by this invention or molecular complex whose structure is unknown more quickly and efficiently than attempting to determine such information ab initio.

Molecular replacement provides an accurate estimation of the phases for an unknown structure. Phases are a factor in equations used to solve crystal structures that can not be determined directly. Obtaining accurate values for the phases, by methods other than molecular replacement, is a time-consuming process that involves iterative cycles of approximations and refinements and greatly hinders the solution of crystal structures. However, when the crystal structure of a protein containing at least a homologous portion has been solved, the phases from the known structure provide a satisfactory estimate of the phases for the unknown structure.

Thus, this method involves generating a preliminary model of a molecule or molecular complex whose structure coordinates are unknown, by orienting and positioning the relevant portion of the AR-LBD/AR-LBD ligand complex according to Table A within the unit cell of the crystal of the unknown molecule or molecular complex so as best to account for the observed X-ray diffraction pattern of the crystal of the molecule or molecular complex whose structure is unknown. Phases can then be calculated from this model and combined with the observed Xray diffraction pattern amplitudes to generate an electron density map of the structure whose coordinates are unknown. This, in turn, can be subjected to any well-known model building and structure refinement techniques to provide a final, accurate structure of the unknown crystallized molecule or molecular complex [E. Lattman, "Use of the Rotation and Translation Functions", in Meth. Enzymol., 115, pp. 55-77 (1985); M. G. Rossmann, ed., "The Molecular Replacement Method", Int. Sci. Rev. Set., No. 13, Gordon & Breach, New York (1972)].

The structure of any portion of any crystallized molecule or molecular complex, or mutant, homologue or orphan receptor that is sufficiently homologous to any portion of the AR-LBD/AR-LBD ligand complex can be solved by this method. Along with the aforementioned AR, there also exist a number of AR for which the activating or

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deactivating ligands may not be characterized. These proteins are classified as AR due to strong sequence homology to other AR, and are known as orphan receptors.

The structure coordinates are also particularly useful to solve the structure of crystals of AR-LBD/AR-LBD ligand co-complexed with a variety of chemical entities. This approach enables the determination of the optimal sites for interaction between chemical entities, including interaction of candidate AR inhibitors with the complex. For example, high resolution X-ray diffraction data collected from crystals exposed to different types of solvent allows the determination of where each type of solvent molecule resides. Small molecules that bind tightly to these sites can then be designed and synthesized and tested for their AR inhibition activity.

All of the complexes referred to above may be studied using well-known X-ray diffraction techniques and may be refined versus 1.5-3 A resolution X-ray data to an R value of about 0.20 or less using computer software, such as X-PLOR [Yale University, 1992, distributed by Molecular Simulations, Inc.; see, e.g., Blundell & Johnson, supra; Meth. Enzymol., vol. 114 & 115, H. W. Wyckoff et al., eds., Academic Press (1985)]. This information may thus be used to optimize known AR agonists, partial agonists, antagonists, partial antagonists and SARMS, and more importantly, to design new AR agonists/antagonists.

Accordingly, the present invention is also directed to a binding site in AR-LBD for an AR-LBD ligand in which a portion of AR-LBD ligand is in van der Walls contact or hydrogen bonding contact with at least one of the following residues: V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906 of AR-LBD. For purposes of this invention, by AR-LBD binding site it is also meant to include mutants or homologues thereof. In a preferred embodiment, the mutants or homologues have at least 25% identity, more preferably 50% identity, more preferably 75% identity, and most preferably 95% identity to residues V685, L700, L701, S702, S703, L704, N705, E706, L707,

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G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906 of AR-LBD binding sites.

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The present invention is also directed to a machine-readable data storage medium, comprising a data storage material encoded with machine readable data, wherein the data is defined by the structure coordinates of an AR-LBD/AR-LBD ligand according to Table A or a homologue of said complex, wherein said homologue comprises backbone atoms that have a root mean square deviation from the backbone atoms of the complex of not more than 3.0Å. Preferably, the machine-readable data storage medium, according to the invention, is wherein said molecule or molecular complex is defined by the set of structure coordinates for AR-LBD/AR-LBD ligand according to Table A, or a homologue of said molecule or molecular complex, said homologue having a root mean square deviation from the backbone atoms of said amino acids of not more than 2.0 Å. In a preferred embodiment the machine-readable data storage medium comprises a data storage material encoded with a first set of machine readable data comprising a Fourier transform of at least a portion of the structural coordinates for an AR-LBD/AR-LBD ligand according to Table A; which, when combined with a second set of machine readable data comprising an X-ray diffraction pattern of a molecule or molecular complex of unknown structure, using a machine programmed with instructions for using said first set of data and said second set of data, can determine at least a portion of the structure coordinates corresponding to the second set of machine readable data, said first set of data and said second set of data.

The present invention also provides for computational methods using three dimensional models of the androgen receptor that are based on crystals of AR-LBD/AR-LBD ligand complex. Generally, the computational method of designing an androgen receptor ligand determines which amino acid or amino acids of the AR-LBD interact with a chemical moiety (at least one) of the ligand using a three dimensional model of a crystallized protein comprising the AR-LBD with a bound ligand, and selecting a chemical modification (at least one) of the

chemical moiety to produce a second chemical moiety with a structure that either decreases or increases an interaction between the interacting amino acid and the second chemical moiety compared to the interaction between the interacting amino acid and the corresponding chemical moiety on the natural hormone.

The computational methods of the present invention are for designing androgen receptor synthetic ligands using such crystal and three dimensional structural information to generate synthetic ligands that modulate the conformational changes of the androgen receptor's LBD. These computational methods are particularly useful in designing an agonist, partial agonist, antagonist or partial antagonist or SARMs to the androgen receptor, wherein the agonist, partial agonist, antagonist or partial antagonist or SARMS has an extended moiety that prevents any one of a number of ligand-induced molecular events that alter the receptor's influence on the regulation of gene expression, such as preventing the normal coordination of the activation domain observed for a naturally occurring ligand or other ligands that mimic the naturally occurring ligand, such as an agonist. As described herein, synthetic ligands of the androgen receptor will be useful in modulating androgen receptor activity in a variety of medical conditions.

AR is known to comprise various domains as follows:

1) a variable amino-terminal domain;

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- 2) a highly conserved DNA-binding domain (DBD); and
- 3) a less conserved carboxyl-terminal ligand-binding domain (LBD).
 25 This modularity permits different domains of each protein to separately accomplish different functions, although the domains can influence each other. The separate function of a domain is usually preserved when a particular domain is isolated from the remainder of the protein. Using conventional protein chemistry techniques a modular domain can sometimes be separated from the parent protein. Using conventional molecular biology techniques each domain can usually be separately expressed with its original function intact or chimerles of two different nuclear receptors can be constructed, wherein the chimetics retain the properties of the individual functional domains of the respective nuclear
 - receptors from which the chimerica were generated.

 Amino Terminal Domain

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The amino terminal domain is the least conserved of the three domains. This domain is involved in transcriptional activation and in some cases its uniqueness may dictate selective receptor-DNA binding and activation of target genes by specific receptor isoforms. This domain can display synergistic and antagonistic interactions with the domains of the LBD. For example, studies with mutated and/or deleted receptors show positive cooperativity of the amino and carboxy terminal domains. In some cases, deletion of either of these domains will abolish the receptor's transcriptional activation functions.

10 DNA-Binding Domain

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The DBD is the most conserved domain. The DBD contains two perpendicularly oriented a-helixes that extend from the base of the first and second zinc fingers. The two zinc fingers function in concert along with non-zinc finger residues to direct nuclear receptors to specific target sites on DNA and to align receptor homodimer or heterodimer interfaces. Various amino acids in DBD influence spacing between two half-sites for receptor dimer binding.

Ligand or AR Binding Domain

The LBD is the second most highly conserved domain. Whereas integrity of several different LBD sub-domains is important for ligand binding, truncated molecules containing only the LBD retain normal ligand-binding activity. This domain also participates in other functions, including dimerization, nuclear translocation and transcriptional activation. Importantly, this domain is the binding site for ligands, i.e. AR modulators, and undergoes ligand-induced conformational changes as detailed herein.

As described herein, the LBD of AR can be expressed, crystallized, its three dimensional structure determined with a ligand bound (either using crystal data from the same receptor or a different receptor or a combination thereof), and computational methods used to design ligands to its LBD, particularly ligands that contain an extension moiety that coordinates the activation domain of AR.

Once a computationally designed ligand (CDL) is synthesized, it can be tested using assays to establish its activity as an agonist, partial agonist, antagonist or partial antagonist or SARM, and affinity, as described herein. After such testing, the CDLs can be further refined by

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generating LBD crystals with a CDL bound to the LBD. The structure of the CDL can then be further refined using the chemical modification methods described herein for three dimensional models to improve the activity or affinity of the CDL and make second generation CDLs with improved properties, such as that of a super agonist or antagonist.

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Typically AR-LBD is purified to homogeneity for crystallization. Purity of AR-LBD is measured with SDS-PAGE, mass spectrometry and hydrophobic HPLC. The purified AR for crystallization should be at least 97.5 % pure or 97.5%, preferably at least 99.0% pure or 99.0% pure, more preferably at least 99.5% pure or 99.5% pure.

Initially purification of the unliganded receptor can be obtained by conventional techniques, such as hydrophobic interaction chromatography (HPLC), ion exchange chromatography (HPLC), and heparin affinity chromatography.

To achieve higher purification for improved crystals of AR, it will be desirable to ligand shift purify the nuclear receptor using a column that separates the receptor according to charge, such as an ion exchange or hydrophobic interaction column, and then bind the eluted receptor with a ligand, especially an agonist or partial agonist. The ligand induces a change in the receptor's surface charge such that when rechromatographed on the same column, the receptor then elutes at the position of the liganded receptor are removed by the original column run with the unliganded receptor. Usually saturating concentrations of ligand are used in the column and the protein can be preincubated with the ligand prior to passing it over the column.

More recently developed methods involve engineering a "tag" such as with histidine placed on the end of the protein, such as on the amino terminus, and then using a nickle chelation column for purification, Janknecht R., Proc. Natl. Acad.Sci. USA Vol 88:8972-8976 (1991) incorporated by reference.

To determine the three dimensional structure of a AR-LBD, it is desirable to co-crystalize the LBD with a corresponding LBD ligand.

Typically purified AR-LBD is equilibrated at a saturating concentration of ligand at a temperature that preserves the integrity of the protein. Ligand equilibration can be established between 2 and 37° C, although the receptor tends to be more stable in the 2-20° C range.

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Preferably crystals are made with the hanging drop methods. Regulated temperature control is desirable to improve crystal stability and quality. Temperatures between 4 and 25°C are generally used and it is often preferable to test crystallization over a range of temperatures. It is preferable to use crystallization temperatures from 18 to 25°C, more preferably 20 to 23°C, and most preferably 22°C.

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Ligands that interact with AR can act as an agonist, partial agonist, antagonist or partial antagonist or SARM based on what ligand-induced conformational changes take place.

Agonists or partial agonists induce changes in receptors that place them in an active conformation that allows them to influence transcription, either positively or negatively. There may be several different ligand-induced changes in the receptor's conformation.

Antagonists or partial antagonists bind to receptors, but fail to induce conformational changes that alter the receptor's transcriptional regulatory properties or physiologically teleram conformations. Binding of an antagonist or partial antagonist can also block the binding and therefore the actions of an agonist or partial agonist.

Partial agonists, or partial antagonists, bind to receptors and induce only part of the changes in the receptors that are induced by agonists or antagonists, respectively. The differences can be qualitative or quantitative. Thus, a partial agonist or partial antagonist may induce some of the conformation changes induced by agonists or antagonists, respectively, but not others, or it may only induce certain changes to a limited extent.

As described herein, the unliganded receptor is in a configuration that is either inactive, has some activity or has repressor activity. Binding of agonist ligands induces conformational changes in the receptor such that the receptor becomes more active, either to stimulate or repress the expression of genes. The receptors may also have non-genomic actions, some of the known types of changes and/or the sequelae of these are listed herein.

Heat shock protein binding domains present a region for binding to the LBD and can be modulated by the binding of a ligand to the LBD. Consequently, an extended chemical moiety (or more) from the ligand that stabilizes the binding or comact of the heat shock protein binding

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domain with the LBD can be designed. Typically such chemical moieties will extend past and away from the molecular recognition domain on the ligand and usually past the buried binding cavity of the ligand.

Ligand binding by the receptor is a dynamic process, which regulates receptor function by inducing an altered conformation.

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The three-dimensional structure of the liganded AR receptor will greatly aid in the development of new AR synthetic ligands. In addition, AR is overall well suited to modern methods including three-dimensional structure elucidation and combinatorial chemistry such as those disclosed in EP 335 628, U.S. patent 5,463,564, which are incorporated herein by reference. Computer programs that use crystallography data when practicing the present invention will enable the rational design of ligand to AR. Programs such as RASMOL can be used with the atomic coordinates from crystals generated by practicing the invention or used to practice the invention by generating three dimensional models and/or determining the structures involved in ligand binding. Computer programs such as INSIGHT and GRASP allow for further manipulation and the ability to introduce new structures. In addition, high throughput binding and bioactivity assays can be devised using purified recombinant protein and modern reporter gene transcription assays described herein and known in the art in order to refine the activity of a CDL.

Generally the computational method of designing an AR synthetic ligand comprises two steps:

- 1) determining which amino acid or amino acids of AR- LBD interacts with a first chemical moiety (at least one) of the ligand using a three dimensional model of a crystallized protein comprising an AR-LBD with a bound ligand; and
- 2) selecting a chemical modifications (at least one) of the first chemical moiety to produce a second chemical moiety with a structure to either decrease or increase an interaction between the interacting amino acid and the second chemical moiety compared to the interaction between the interacting amino acid and the first chemical moiety.
- Preferably the method is carried out wherein said three dimensional model is generated by comparing isomorphous ligand derivatives to produce improved phasing. Further preferred is wherein said method

comprises determining a change in interaction between said interacting amino acid and said ligand after chemical modification of said first chemical moiety, especially wherein said three dimensional model is generated by comparing isomorphous ligand derivatives to produce improved phasing. Also preferred is wherein said selecting uses said first chemical moiety that interacts with at least one of the interacting amino acids V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906.

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As shown herein, interacting amino acids form contacts with the ligand and the center of the atoms of the interacting amino acids are usually 2 to 4 angstroms away from the center of the atoms of the ligand. Generally these distances are determined by computer as discussed herein and in McRee 1993, however distances can be determined manually once the three dimensional model is made. See also Wagner et al., Nature 378(6558):670-697 (1995) for stereochemical figures of -three dimensional models. More commonly, the atoms of the ligand and the atoms of interacting amino acids are 3 to 4 angstroms apart. The invention can be practiced by repeating steps I and 2 to refine the fit of the ligand to the LBD and to determine a better ligand, such as an agonist, partial agonist, antagonist or partial antagonist or SARM. The three dimensional model of AR can be represented in two dimensions to determine which amino acids contact the ligand and to select a position on the ligand for chemical modification and changing the interaction with a particular amino acid compared to that before chemical modification. The chemical modification may be made using a computer, manually using a two dimensional representation of the three dimensional model or by chemically synthesizing the ligand. The ligand can also interact with distant amino acids after chemical modification of the ligand to create a new ligand. Distant amino acids are generally not in contact with the ligand before chemical modification. A chemical modification can change the structure of the ligand to make as new ligand that interacts with a distant amino acid usually at least 4.5

angstroms away from the ligand, preferably wherein said first chemical moiety is 6 to 12 angstroms away from a distant amino acid. Often distant amino acids will not line the surface of the binding cavity for the ligand, they are too far away from the ligand to be part of a pocket or binding cavity. The interaction between a LBD amino acid and an atom 5 of an LBD ligand can be made by any force or attraction described in nature. Usually the interaction between the atom of the amino acid and the ligand will be the result of a hydrogen bonding interaction, charge interaction, hydrophobic interaction, van der Waals interaction or dipole interaction. In the case of the hydrophobic interaction it is recognized 10 that this is not a per se interaction between the amino acid and ligand, but rather the usual result, in part, of the repulsion of water or other hydrophilic group from a hydrophobic surface. Reducing or enhancing the interaction of the LBD and a ligand can be measured by calculating or testing binding energies, computationally or using thermodynamic or 15 kinetic methods as known in the art.

Chemical modifications will often enhance or reduce interactions of an atom of a LBD amino acid and an atom of an LBD ligand. Steric hindrance will be a common means of changing the interaction of the LBD binding cavity with the activation domain.

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The present invention also provides methods for identifying compounds that modulate androgen receptor activity. Various methods or combinations thereof can be used to identify these compounds. For example, test compounds can be modeled that fit spatially into the AR-LBD as defined by structure coordinates according to Table A, or using a 25 three-dimensional structural model of AR-LBD, mutant AR-LBD or AR-LBD homolog or portion thereof. Structure coordinates of the ligand binding site, in particular amino acids V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, 30 R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906 can also be used to identify structural and chemical features. Identified structural or chemical features can then be employed to design or select 35 compounds as potential AR modulators. By structural and chemical

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features it is meant to include, but is not limited to, van der Waals interactions, hydrogen bonding interactions, charge interaction, hydrophobic bonding interaction, hydrophobic interaction and dipole interaction. Alternatively, or in conjunction, the three-dimensional structural model or the ligand binding site can be employed to design or 5 select compounds as potential AR modulators. Compounds identified as potential AR modulators can then be synthesized and screened in an assay characterized by binding of a test compound to the AR-LBD. Examples of assays useful in screening of potential AR modulators include, but are not limited to, screening in silico, in vitro assays and 10 high throughput assays. Finally, these methods may also involve modifying or replacing one or more amino acids from AR-LBD such as V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, 15 M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906 of AR-LBD according to Table A.

A preferred method of the invention can be described as a computational method of designing an androgen receptor antagonist from an androgen receptor agonist comprising:

- 1) determining a structure of a molecular recognition domain of said agonist using a three dimensional model of a crystallized protein comprising an AR-LBD, and
- 2) selecting at least one chemical modification of said agonist that provides a ligand structure that extends beyond a binding site for said agonist and in the direction of at least one protein domain important in AR biological function.

Another preferred method of the invention can be described as a computational method of designing a selective androgen receptor modulator such as an androgen receptor super agonist or antagonist comprising:

 determining at least one interacting amino acid of an AR-LBD that interacts with at least one first chemical moiety of said ligand using a three dimensional model of a crystallized protein comprising AR-LBD with a bound ligand, and

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2) selecting at least one chemical modification of said first chemical moiety to produce a second chemical moiety with a structure to reduce or enhance an interaction between said interacting amino acid and said second chemical moiety compared to said interaction between said interacting amino acid and said first chemical moiety.

However, as will be understood by those of skill in the art upon this disclosure, other structure based design methods can be used. Various computational structure based design methods have been disclosed in the art.

For example, a number computer modeling systems are available in which the sequence of the AR-LBD and the AR-LBD structure (i.e., atomic coordinates of AR-LBD and/or the atomic coordinates of the active site, the bond and dihedral angles, and distances between atoms in the active site such as provided in Table A) can be input. This computer system then generates the structural details of the site in which a potential AR modulator binds so that complementary structural details of the potential modulators can be determined. Design in these modeling systems is generally based upon the compound being capable of physically and structurally associating with AR-LBD. In addition, the compound must be able to assume a conformation that allows it to associate with AR-LBD. Some modeling systems estimate the potential inhibitory or binding effect of a potential AR modulator prior to actual synthesis and testing.

Methods for screening chemical entities or fragments for their ability to associate with AR-LBD are also well known. Often these methods begin by visual inspection of the active site on the computer screen. Selected fragments or chemical entities are then positioned with the AR-LBD. Docking is accomplished using software such as QUANTA and SYBYL, following by energy minimization and molecular dynamics with standard molecular mechanic forcefields such as CHARMM and AMBER. Examples of computer programs which assist in the selection of chemical fragment or chemical entities useful in the present invention include, but are not limited to, GRID (Goodford, P.J. J. Med. Chem. 1985 28:849-857), AUTODOCK (Goodsell, D.S. and Olsen, A.J. Proteins,

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Structure, Functions, and Genetics 1990 8:195-202), and DOCK (Kunts et al. J. Mol. Biol. 1982 161:269-288).

Upon selection of preferred chemical entities or fragments, their relationship to each other and AR-ABD can be visualized and the entities or fragments can be assembled into a single potential modulator. Programs useful in assembling the individual chemical entities include, but are not limited to CAVEAT (Bartlett et al. Molecular Recognition in Chemical and Biological Problems Special Publication, Royal Chem. Soc. 78, 182-196 (1989)) and 3D Database systems (Martin, Y.C. J. Med. Chem. 1992 35:2145-2154).

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Alternatively, compounds may be designed *de novo* using either an empty active site or optionally including some portion of a known inhibitor. Methods of this type of design include, but are not limited to LUDI (Bohm H-J, J. Comp. Aid. Molec. Design 1992 6:61-78) and LeapFrog (Tripos Associates, St. Louis. MO).

The present invention is also directed to an AR-LBD selective androgen receptor modulator (SARM), in particular an agonist or antagonist or partial agonist or partial antagonist, identified by a computational process of the invention.

The present invention is further directed to a method for treating prostate cancer comprising administering an effective amount of an AR modulator, preferably an antagonist or partial antagonist, identified by a computational process of the invention.

The present invention is also direct to a method for treating an age related disease comprising administering an effective amount of an AR modulator, preferably an agonist or partial agonist, identified by a computational process of the invention, preferably wherein said age related disease is osteoporosis, muscle wasting or loss of libido.

Compounds identified as agonists, partial agonists, antagonists, partial antagonists or SARMs by the methods disclosed herein which are active when given orally can be formulated as liquids for example syrups, suspensions or emulsions, tablets, capsules and lozenges. A liquid composition will generally consist of a suspension or solution of the compound in a suitable liquid carrier(s), for example ethanol, glycerin, sorbitol, non-aqueous solvent such as polyethylene glycol, oils or water, with a suspending agent, preservative, surfactant, wetting agent,

flavoring or coloring agent. Alternatively, a liquid formulation can be prepared from a reconstitutable powder. For example a powder containing active compound, suspending agent, sucrose and a sweetener can be reconstituted with water to form a suspension; and a syrup can be prepared from a powder containing active ingredient, sucrose and a 5 sweetener. A composition in the form of a tablet can be prepared using any suitable pharmaceutical carrier(s) routinely used for preparing solid compositions. Examples of such carriers include magnesium stearate, starch, lactose, sucrose, microcrystalline cellulose, binders, for example polyvinylpyrrolidone. The tablet can also be provided with a color film 10 coating, or color included as part of the carrier(s). In addition, active compound can be formulated in a controlled release dosage form as a tablet comprising a hydrophilic or hydrophobic matrix. A composition in the form of a capsule can be prepared using routine encapsulation procedures, for example by incorporation of active compound and 15 excipients into a hard gelatin capsule. Alternatively, a semi-solid matrix of active compound and high molecular weight polyethylene glycol can be prepared and filled into a hard gelatin capsule; or a solution of active compound in polyethylene glycol or a suspension in edible oil, for example liquid paraffin or fractionated coconut oil can be prepared and 20 filled into a soft gelatin capsule. Compounds identified by the processes described herein which are active when given parenterally can be formulated for intramuscular or intravenous administration. A typical composition for intra-muscular administration will consist of a suspension or solution of active ingredient in an oil, for example arachis 25 oil or sesame oil. A typical composition for intravenous administration will consist of a sterile isotonic aqueous solution containing, for example active ingredient, dextrose, sodium chloride, a co-solvent, for example polyethylene glycol and, optionally, a chelating agent, for example ethylenediaminetetracetic acid and an anti-oxidant, for example, sodium 30 metabisulphite. Alternatively, the solution can be freeze dried and then reconstituted with a suitable solvent just prior to administration. Identified compounds which are active on rectal administration can be formulated as suppositories. A typical suppository formulation will generally consist of active ingredient with a binding 35 and/or lubricating agent such as a gelatin or cocoa butter or other low

melting vegetable or synthetic wax or fat. Identified compounds which are active on topical administration can be formulated as transdermal compositions. Such compositions include, for example, a backing, active compound reservoir, a control membrane, liner and contact adhesive.

The typical daily dose of a varies according to individual needs, the condition to be treated and with the route of administration. Suitable doses are in the general range of from 0.001 to 10 mg/kg bodyweight of the recipient per day.

The following examples are to illustrate the invention, but should not be interpreted as a limitation thereon.

Examples

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Cloning, Expression and Purification of the Androgen Receptor Ligand-Binding Domain

The rat androgen receptor (rAR) ligand-binding domain (LBD) cDNA, from amino acid 646 to 901, was cloned from a rat prostate cDNA 15 library (Clontech) by PCR. The primers used were CATATGATTGAAGGCTATGAATGTCAACCTATCTTT (SEQ ID NO:3) and TCACTGTGTGTGGAAATAGATGGG (SEQ ID NO:4). The rat AR LBD was expressed as a fusion protein driven by the T7 promoter of pET28b vector (Novagen) to include an N-terminal polyhistidine tag and a 20 thrombin cleavage site. The replacement of T877 for A (the LNCaP mutation) in this rAR LBD expression construct was performed with the QuickChange Site-Directed Mutagenesis kit (STRATAGENE). Dihydrotestosterone (DHT) was included in the E. coli (BL21-DE3) fermentation medium at a concentration of 0.05 mM. Induction with 25 $0.4\ mM$ isopropyl- β -D-thiogalactopyranoside was allowed to proceed for 16 hours at 20°C in M9 minimal media supplemented with casamino acids (Difco) and trace minerals, and pellets were stored at -70 °C. A total of 6-9 mg of recombinant AR LBD was isolated from a 15 gram cell pellet following sonication and chromatography on a nickel-chelate resin. 30 Polyhistidine-tagged AR LBD of approximately 90% purity eluted at 0.45 M imidazole in a gradient of 0.05-1.0 imidazole. This material was quantitatively cleaved at an engineered site for thrombin recognition, followed by chromatography on benzamidine sepharose (Pharmacia) to remove the serine protease, with a 70% recovery. The final sample 35 containing the sequence Gly-Ser-His-Met at the N-terminus followed by

residues 646-901 of the rat (664 – 919 in the human) AR LBD protein, was concentrated for crystallography to 2 mg/ml in 20 mM Tris (pH 7.5), 0.5 M NaCl, 10% glycerol, 1 mM EDTA and 1 mM DTT.

The sequence of the rat Androgen Receptor LBD (AR), as cloned, with the secondary structural features marked. For comparison, the aligned sequence of the Progesterone Receptor LBD (PR) is given. Residues involved in androgen binding are marked (*). Residues which are disordered in the crystal structure are underlined. The AR sequence is SEQ ID NO: 1. The PR sequence is SEQ ID NO: 2.

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                           |-----H3-----
     |-H1--|
    660 GSHMIEGYECQPIFLNVLEAIEPGVVCAGHDNNQPDSFAALLSSLNELGE
                                                           AR
            GQDIQLIPPLINLLMSIEPDVIYAGHDNTKPDTSSSLLTSLNQLGE
    678
15
                            |-----H4/5-----
     710 RQLVHVVKWAKALPGFRNLHVDDQMAVIQYSWMGLMVFAMGWRSFTNVNS
                                                           AR
    724 RQLLSVVKWSKSLPGFRNLHIDDQITLIQYSWMSLMVFGLGWRSYKHVSG
20
                               |----H7----
          SSSS SSS |-H6|
     760 RMLYFAPDLVFNEYRMHKSRMYSQCVRMRHLSQEFGWLQITPQEFLCMKA
     774 QMLYFAPDLILNEQRMKESSFYSLCLTMWQIPQEFVKLQVSQEEFLCMKV
                        |----H9-----
25
     810 LLLFSIIPVDGLKNOKFFDELRMNYIKELDRIIACKRKNPTSCSRRFYQL
                                                           AR
     824 LLLLNTIPLEGLRSQTQFEEMRSSYIRELIKAIGLRQKGVVSSSQRFYQL
                                        |-----H12-----|
         ---H10/11-----| |--|
     860 TKLLDSVQPIARELHQFTFDLLIKSHMVSVDFPEMMAEIISVQVPKILSG
                                                           AR
30
     874 TKLLDNLHDLVKQLHLYCLNTFIQSRALSVEFPEMMSEVIAAQLPKILAG
          SSS
     910 KVKPIYFHTQ
                    AR
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     924 MVKPLLFHK
                    PR
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Crystallization

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The AR-LBD - Dihydrotestosterone (DHT) complex was crystallized at 20° C by vapor diffusion in the hanging-drop mode. In the crystallization trials, the protein complex as obtained from MMB&B was used without any further purification. In the initial trial to obtain crystallization conditions, a sparse matrix crystallization screen was done with the Crystal Screens 1 and 2 (Hampton Research). For each crystallization trial, a 2 µl drop was prepared by mixing 1 µl of purified protein (1.9 mg ml⁻¹) with an equal volume of reservoir solution. The reservoir contained 1.0 ml of the precipitating solution. Small crystals were obtained in two days from six of the drops (table 1).

Table 1: Crystallization Conditions

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# Precipitating Solution	Result
1.5 M Li Sulfate, 0.1M Na Hepes, pH 7.5	Small r ds
0.8 M Na/K Tartrate, 0.1M Na Hepes, pH 7.5	Larger rods
2% v/v PEG 400, 2.0 M Am Sulfate,	
0.1M Na Hepes, pH 7.5	Small cubes
1.6 M Mg Sulfate, 0.1M MES, pH 6.5	Small crystallites
1.6 M Am Sulfate, 0.1 M Na Cl,	
0.1 M Hepes, pH 7.5	Small rods
12% v/v Glycerol, 1.5 M Am Sulfate,	
0.1 M Tris, pH 8.5	Small rods
	0.8 M Na/K Tartrate, 0.1M Na Hepes, pH 7.5 2% v/v PEG 400, 2.0 M Am Sulfate, 0.1M Na Hepes, pH 7.5 1.6 M Mg Sulfate, 0.1M MES, pH 6.5 1.6 M Am Sulfate, 0.1 M Na Cl, 0.1 M Hepes, pH 7.5 12% v/v Glycerol, 1.5 M Am Sulfate,

The largest single crystal, measuring 0.05 mm x 0.04 mm x 0.26mm, was obtained from Crystal Screen 1, solution # 29 (0.8 M Na/K Tartrate, 0.1M Na Hepes, pH 7.5). This crystal was subsequently used in the initial data collection run (as described below).

Optimization of the crystallization condition was done using a Cyperlab C-200 automated crystallization robotic workstation. A crystallization trial was performed using a 24-step linear gradient from 0.6 M to 1.26 M Na tartrate, 100 Mm Hepes, pH 7.5 (Note: The optimization screen used sodium rather than sodium/potassium tartrate). The largest, rod shaped crystal, with dimensions 0.09 mm x 0.09 mm x 0.20mm, was obtained at 0.887 M Na Tartrate. This crystal was used in the second data collection run (as described below).

Data Collection and Reduction

For the initial X-ray experiment, the crystal from the initial crystallization screen was flash cooled by dipping it in a cryoprotectant solution containing the precipitating solution (0.8 M Na/K Tartrate, 0.1M Na Hepes, pH 7.5) with 250mm NaCl and 20% Glycerol added and then placed it in a cold stream at 100° K.

For data set 1, X-ray diffraction data were collected with an R-Axis II imaging plate detector. The radiation was generated from a Rigaku RU-200 rotating at 5 kw power with a fine focus filament (0.3 x 3.0mm) was monchromated (Cu Kα) and intensified by focusing with Yale mirrors (Molecular Structure Corporation). The crystal diffracted to better than 2.4 Å resolution. Autoindexing and processing of the measured intensity data was carried out with the HKL software package (Otwinoski, L. (1993) in CCP4 Study Weekend, Data Collection and

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Processing (Sawyer, L., Issacs, N., and Bailey, S., Eds.) pp 56-62, SERC Daresbury Laboratory, Warrington, U.K). X-ray diffraction from the crystals have the symmetry and systematic absences of the orthorhombic space group P212121 with unit cell dimensions a = 56.03 Å, b = 66.27 Å, c = 70.38 Å, and one molecule per asymmetric unit (Mathews Volume = 2.16 Å 3 Da-1).

A second X-ray diffraction data set (data set 2) was collected at the IMCA-CAT beamline (sector 17ID) at the Advanced Photon Source synchrotron at Argonne, II. The crystal from the optimization screen described above, was flash-cooled by placing it in the reservoir solution (0.877 M Na Tartrate, 0.1M Na Hepes, pH 7.5) with 250mm NaCl and 20% Glycerol added, and then placing it in a cold stream at 100° K. The data were collected with a Bruker 2x2 mosaic CCD detector. The crystal diffracted to better than 2.0 Å. Autoindexing and processing of the measured intensity data was carried out with the HKL2000 software package (Otwinoski, L. (1993) in CCP4 Study Weekend, Data Collection and Processing (Sawyer, L., Issacs, N., and Bailey, S., Eds.) pp 56-62, SERC Daresbury Laboratory, Warrington, U.K.). The data collection and processing statistics for both data sets are summarized in table 2.

Structure Determination (Molecular Replacement)

The structure was determined by the method of molecular replacement with the program AmoRe (Navaza, J. (1994) AmoRe: an automated package for molecular replacement. Acta Cryst. D50, 157-163). The Progesterone Receptor ligand binding domain (PR-LBD), which has 54% sequence identity and 76% sequence homology to AR-LBD, was used as the search model. The atomic coordinates of PR-LBD (Protein Data Bank reference code 1A28) by Williams & Sigler (Nature 1998 393, 391) were unmodified except for the removal of the ligand and solvent molecules. A second molecular replacement search was performed with a theoretical model for the AR-LBD provided by the MMS/CADD group (table 3). The PR-LBD structure gave a slightly better solution than the AR-model (1.7 σ vs.1.3 σ above background) and was used in the subsequent refinement, although both structures gave equivalent results with no molecular interpenetration.

Table 2:	Data C	11	cti	n	and	Proc	ssing
						Data C	nt 1

	Table 2: Data C II cti n and	Data Set I	Data Set II
	Date	5/19/99	6/17/99
	Source/Detector	Rigaku RU-200	IMCA/APS 17ID
5	Detector	R-axis II	Bruker 2x2
	Wavelength	Cu Kα (1.54 Å)	1.00 Å
	Frames	364	400
	ΔΦ	0.5°	0.5°
	Crystal to plate distance	150 mm	135 mm
10	Time/frame	20 min	1 sec
	Number of Observations	209,891	416,207
	Data Reduction Program	HKL	HKL2000
	Unique reflections	10,824	18,308
	Reflections Used	10,114	16,862
15	Resolution	2.4 Å (2.5-2.4 Å)	2.0 Å (2.1-2.0 Å)
	Completeness	93.8% (71.6%)	92.6 % (73.0 %)
	Multiplicity	6.3	7.3
	Mosiacity	0.502	0.332
	Rsym (on I)	4.2 % (17.5%)	10.1 % (25.6%)
20	Space Group	P212121	P212121
	a	56.09 Å	56.08 Å
	b	66.43 Å	65.76 Å
	c	70.54 Å	70.51 Å
	Wilson B-value	39.05 Ų	29.26 Ų

Values for data in the last resolution shell are given in parentheses

Table 3: Molecular Replacement Statistics

	Search Model:	Progesterone	AR Model
30		(PDB file 1A28)	
	Program Used	AmoRe	AMoRe
	Resolution Range	8.0 – 4.0 Å	8.0 – 4.0 Å
	Radius of Integration	25 Å	25 Å
	Number of Reflections	2.393	2,393
35	Number of Atoms	2,019	2,094
	RF Correlation (2 nd solution)	0.16 (0.12)	0.13 (0.11)
	TF Correlation (2 nd solution)	0.31 (0.20)	0.23 (0.14)
	TF R-factor (2 nd solution)	49.0% (52.7%)	52.1% (54.0%)
	Rigid Body Correlation	0.34	0.28
40	Rigid Body R-factor	48.1%	50.4%

Structure Refinement

The structure was first refined with the initial 2.4 Å data set (2σ) data, 9,818 reflections) by the method of simulated annealing with

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program X-PLOR (Brünger, A.T., Kuriyan, J. & Karplus, J. (1987) "Crystallographic R-factor refinement by molecular dynamics", Science 235: 458-460) in four cycles to an R-factor of 27.7%. Each refinement cycle consisted of a least-squares minimization, simulated annealing at 3000°, and individual isotropic B-factor refinement. The first cycle, with the Progesterone molecular replacement model unmodified for the sequence differences between AR and PR, gave an R-factor of 33.8%. The model was then rebuilt using the AR amino acid sequence and a second refinement cycle gave an R-factor of 29.6%. At this stage of the refinement, the DHT molecule could be clearly seen in the difference electron density map.

After each cycle, the structure was carefully examined using molecular computer graphics program Chain (Sack, John S. (1988) "CHAIN- A Crystallographic Modeling Program", J. Mol. Graphics 6: 224-225) and modifications were made to the structure as needed. Several 15 residues, from both the N- and C-termini of the molecule, which were not seen in the electron density maps were removed from the model. After the second cycle of refinement, the DHT was added to the model. Solvent molecules were added where there were 3σ peaks in both the 2Fo - Fcand Fo - Fc electron density maps and removed if their B-factor went 20 above 60 Å². After four cycles of X-PLOR refinement, a careful examination of the electron density showed the model to be much improved, although molecular refitting still needed to be done in some regions. The density is clear except for some of the loop regions, particularly the loop between helices I and II, which was also poorly 25 modeled in the PR structure.

Table 4: Refinement Statistics (X-PLOR)

30	Part I: 2σ data (9,818 reflections) to 2.4 Å	
JU	rait i. 20 data (5,0 to reflections) to 2.474	

	•,•.•.			
Cycle 1	251 residues	No ligand	0 waters	R = 33.8 %
Cycle 2	248 residues	No ligand	0 waters	R = 29.6%
Cycle 3	247 residues	ligand	18 waters	R = 28.3 %
Cycle 4	246 residues	ligand	40 waters	R = 27.7%

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Part II: 2o data (15,067 reflections) to 2.0 Å

Cycle 5	246 residues	ligand	32 waters	R = 27.9 %
Cycle 6	246 residues	ligand	57 waters	R = 26.8 %

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 Cycle 7
 246 residues
 ligand
 58 waters
 R = 26.7 %

 Cycle 8
 246 residues
 ligand
 106 waters
 R = 24.2%

At this stage of the refinement, the higher resolution data collected at the APS synchrotron became available. Four additional X-PLOR refinement cycles were performed with the 2.0 Å data set (2 σ data, 15,067 reflections) following the same protocol. The final structure has an R-factor of 24.2% with a total of 106 solvent molecules. The final refinement statistics are presented in table 5.

10 Table 5: Final Refinement Parameters

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10.0 - 2.0 ÅResolution Range 15,067 Reflections 24.2 % R-factor 31.2 % R-free 246 (672-917) 15 # residues 2118 (1991 atoms, 21 DHT, 106 waters) # atoms **RMS** deviations 0.014 Å bond lengths 1.594° bind angles 1.558° 20 Improper angles Average B-factors 25.02 Å² Protein 14.40 Å² DHT 30.21 Å² Water 29.26 Å² Wilson B-factor 25

Description of the Molecule

The structure of AR-LBD is complete from residues 671 through 917 for the wild-type and 672 to 918 for the LNCaP mutant. Analysis of the structures with program PROCHECK showed only minor exceptions to the allowed geometry. In the wild-type structure, the first six residues of the chain (664 - 670) are not seen in the electron density and are probably disordered. This leaves only one residue before the initial residue of the first α-helix (H1) in the wild-type structure, none in the LNCaP mutant structure. On the C-terminal end, the last two residues (918 - 919) are not seen in the electron density of the wild-type structure, but only the last is missing in the mutant. In addition, since the loop between helices 9 and 10 (residues 845-850) is not well defined, it has been modeled as poly-alanine.

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Folding and Packing

As expected, the AR LBD has the same overall three-dimensional structure as those of the other nuclear hormone receptor LBDs. The molecule is folded into a "helical sandwich" consisting of 10 α -helices. There are four small pieces of beta strand, forming two short beta-sheets; one in the core of the molecule between helices 5 and 6 near the ligand binding site, and the other formed by the loop between helices 8 and 9 and the C-terminus. This latter sheet, also seen in the PR LBD structure, holds helix 12 in the closed, agonist conformation, close to and capping the ligand binding site.

Lack of Dimer Formation

Studies have indicated that the estrogen, progesterone, and androgen receptors all function as homodimers and that AR LBD forms dimers in solution. Thus it could be expected that the AR LBD domains 15 might form homodimers in the crystal similar to those previously seen in the RXR- α and estrogen receptor (ER) LBD crystal structures. In the PR LBD structure, the two monomers in the asymmetric unit are related by a dyad, but the two-fold-symmetric configuration is strikingly different from that of the RXR and ER homodimers and the area buried in this 20 configuration is much smaller than would be expected for stable dimer formation. In the AR LBD crystal, the ligand-binding domains are unmistakably monomeric, and there are no twofold axes relating domains. Moreover, the homodimer interaction seen in the structures of ER and RXR LBDs is not possible for the AR LBD, as the C-terminal tail 25 is bound to the groove formed by helices 9 and 10, thereby obstructing the contact region between monomers in RXR and ER homodimers. Whether this observation reflects a non-dimeric state of the AR LBD in the functional AR dimer or is an artifact of the conditions used for AR LBD crystallization remains to be determined. It is noteworthy that the 30 ER LBD constructs used for crystallization have been truncated to remove an analogous C-terminal extension.

Comparison with Progesterone Receptor

While there is only 55% sequence identity between AR LBD and PR LBD, there is a 77% sequence similarity, and as expected, the three-dimensional structures of these two LBDs are very similar with an r.m.s.

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deviation of 1.3 Å between corresponding $C\alpha$ atom positions. As with PR, AR LBD has no helix 2, but its helix 12 is longer than those of RXR or TR. In the case of AR, while helices 10 and 11 are nearly contiguous, there is a proline residue at position 868 that causes a kink between the two helices.

Comparison with theoretical AR model

The theoretical AR model obtained from MMS/CADD and the AR structure have an r.m.s. deviation of 1.29 Å for the 247 alpha carbons. More importantly, the hormone binding site is virtually identical with the exception of the side chains of Met 732(749), Leu 863(880), and Leu 864 (881) which are in different rotomers. This causes the binding cavity to be more compact in the AR structure. Also, there is a flip of the side chain of Asn 688(705) so that the ND2 atom is in position to make a hydrogen bond to the carbonyl off of the D-ring.

15 Table 6: Comparison of AR-LBD to PR-LBD and Theoretical model

	Calpha	Main	Side	IOlai
AR vs. Pr	1.22 (246)	1.27 (983)	1.80 (772)	1.53 (1,755)
AR vs. CADD	1.25 (246)	1.31 (983)	2.41(971)	1.93 (1,954)

20 Binding of Dihydrotestosterone

At the end of the molecular replacement procedure with the PR LBD structure without progesterone as search model, the largest piece of difference electron density, at approximately the 3o level, was found at the progesterone-binding site. Replacing the bound progesterone agonist (which has a carboxyl group at the 17-position) with a model of dhydrotestosterone (DHT, which has a hydroxyl group at the 17-position) produced an even better fit to the difference electron density, indicating that DHT binds to AR LBD in an almost identical fashion to the way progesterone binds to PR LBD. Both agonists interact with helices 3, 5, and 11 of their respective LBDs. Ring A, which is identical in the two steroids, makes similar interactions with the side chains of Q711, M745, R752 (Q725, M759, R766 in PR LBD), and a conserved water molecule. The interactions with ring C are also similar, with close contacts to the mainchain of L704 (L718 in PR LBD) and sidechain of N705 (N719 in PR LBD). The contact between C18 and the O_γ1 of T877 is unique to the wild-type AR LBD, as the corresponding cysteinyl side chain is pointed away from the steroid in the PR LBD structure.

Since progesterone and DHT differ in the substituent on ring D, it is expected that interactions with respective receptors will differ in this region. In the AR LBD structure, No2 of N705 makes a hydrogen bond to the D-ring hydroxyl of DHT. A similar interaction could be made between progesterone and the PR LBD if there were a flip of both the 5 steroid acetyl group and the side chain of N719. This would place the oxygen approximately 3.2 Å from the Nδ2 atom of Asn 719. The ligand contact surface area is slightly larger for progesterone in PR than for DHT in AR (483 vs. 448 Å²) but they are both considerably smaller than the ligand contact surface area in TR (559 Ų), PPAR γ (583 Ų), or the 10

Vitamin D receptor (677 $Å^2$). Figure 3 shows two orthogonal views of the omit electron density map of dihydrotestosterone (DHT) in the hormone-binding site of AR-LBD. There are hydrogen bonds between the steroid and the side chains of Arg 752 and Asn 705.

Table 7: Dihydrotestosterone Contacts (3.4 Å) Hydrogen Bonds

	,		
20	O3 O3	Arg 752 Nh2 Gln 711 Nε2	2.89 Å (2.77 A) 3.36 Å (3.20 A)
25	O20 O20	Asn 705 Nδ2 Thr 877 Ογ1	2.80 Å (3.20 A) 2.70 Å (N/A)
25	Possible Close	e Contacts	
	C11	Leu 704 O	3.31 Å
30	C12	Asn 705 Nδ2	3.07 Å
	C17	Asn 705 Nδ2	3.34 Å
	C19	Met 745 Sδ	3.38 Å
35	C18	Thr 877 Ογ1	3.07 Å

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Comparison with Progesterone binding

Comparison of the structure of DHT in the AR-LBD with the structure of progesterone in the PR-LBD (Williams, S.P. & Sigler, P.B. 40 (1998) "Atomic Structure of Progesterone Complexed with its Receptor", Nature 393, 391) shows a similar mode of binding. Ring A, which is identical in the two steroids, makes similar interactions with the side chains of Q711, M745, R752, Q711 and a conserved water molecule (table 8). The interaction with ring C are also similar, with close 45

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contacts to the mainchain of L704 and sidechain of N705. The contact from C18 to the O γ 1 of T877 is unique to AR-LBD, as the corresponding cysteine sidechain is pointed away from the steroid in the PR-LBD structure

Since progesterone and DHT differ in the substitution off of ring D, it is expected that there will be different interactions with the protein in this region. In the AR structure, the N δ 2 atom of Asn 705 makes hydrogen bond to the D-ring hydroxyl.

A similar interaction could be made in the PR if there were a flip of both the steroid carboxyl group and the side chain of N719. This would place the carboxyl oxygen approximately 3.2 A from the N δ 2 atom of Asn 719. In AR-LBD, there is also a close contact to the side chain of T877 which is absent in the PR-LBD structure.

Figure 4 shows comparison of AR and PR steroid binding Comparison of the binding of dihydrotestosterone to AR-LBD (top) and of progesterone to PR-LBD (bottom). Note that an additional hydrogen bond interaction would be possible if both the sidechains of both N719 and the progesterone were flipped.

Table 8: Comparison of AR and PR steroid binding

20	Table 8. comparison of 12th and 15th										
20		AR	PR .								
	Ring A										
0.5	O3:	H-bond to R752 NH2 (2.9 A)	H-bond to R766 NH2 (2.8 A)								
25		H-bond to water (3.5 A)	H-bond to water (3.1 / 3.4 A)								
		SC of Q711 in different rotomer distance to O3 is 3.4 and 4.13 A	Contact to SC of Gln 725 distance to O3 is 3.2 and 3.3 A								
30	C19	Contact to M745 SD (3.4 A)	Similar orientation (3.5 A)								
35	C2 :	SC of Q711 (3.5 A)	different rotomer (3.2 & 3.3) distance to C4 is 4.1 A								
	Ring C										
	C11	LO704 O (3.3A)	(3.5A)								
40	C12	Contact to N705 Nδ2 (3.1A)	Contact to N719 O81 (3.4 A)								
	C18	Contact T877 Ογ1 (3.1 A)	SC of C891 pointing away distance to Sγ is 3.8 A								
	Ring D										
45	O20/C21	O21 in AR is close to C21 in PR (Pos	ssible flip of Carboxyl in PR?)								

N/A

O20: Contact to C891 Ca (3.2 A)

O20: H-bond N705 Nδ2 (2.8A) O20: Contact T877 Cγ1 (2.7 A) C21: Contact to N719 OD1 (3.2 A) SC of C891 pointing away

C17

Contact N705 Nδ2 (3.3 A)

Ring in slightly different orientation; distance to N719 O81 is 4.7 A

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Structure of the Complex of DHT with the LBD of the LNCaP Mutant

In the LNCaP mutant, T877 is replaced by an alanine residue. The mutant LBD structure has an r.m.s. deviation of 0.8 Å compared to the wild-type structure, close to the expected r.m.s. deviation due to the estimated errors in the coordinates. In particular, the binding of DHT is 15 essentially identical by wild-type and mutant LBDs except at the point of mutation. Here the replacement of T877 by alanine leaves additional space off the D-ring of DHT to accommodate a larger substituent on position 17. This may explain the promiscuous ability of the LNCaP mutant, unlike wild-type AR, to bind to a variety of other hormones and 20 analogs like some progestins, estrogens and cortisols that differ from DHT in substitution at position 17. For example, the binding of flutamide, estradiol, and progesterone to the LNCaP mutant can activate the mutant receptor. Conversely, mutation of T877 to residues with larger sidechains such as aspartic acid and lysine would be expected 25 completely preclude the binding of ligands with any substituent at position 17 of the D-ring and such mutations have been shown to totally eliminate androgen binding.

Table A

	ттом	1	СВ	ILE	672	14.846	25.527	23.734	1.00 25.78
5	MOTA	2		ILE	672	16.247	25.008	24.099	1.00 25.56
3	ATOM	3		ILE	672	14.842	27.035	23.978	1.00 25.60
	MOTA			ILE	672	15.312	27.404	25.360	1.00 25.81
	ATOM	4			672	15.115	23.900	21.789	1.00 25.32
	ATOM	5		ILE		16.189	23.926	21.195	1.00 24.67
	ATOM	6		ILE	672	13.004	25.282	22.008	1.00 24.75
10	ATOM	7		ILE	672			22.242	1.00 25.11
	ATOM	8		ILE	672	14.475	25.215		1.00 25.89
	ATOM	9	N	PHE	673	14.448	22.768	22.030	
	ATOM	10	CA	PHE	673	14.980	21.446	21.635	1.00 25.86
	ATOM	11	CB	PHE	673	14.020	20.306	22.029	1.00 26.22
15	ATOM	12		PHE	673	14.557	18.923	21.722	1.00 25.12
10	ATOM	13	CD1		673	15.765	18.501	22.251	1.00 25.16
		14	CD2		673	13.877	18.066	20.874	1.00 25.81
	ATOM		CEI		673	16.286	17.255	21.946	1.00 23.42
	ATOM	15	CE2		673	14.403	16.809	20.567	1.00 25.08
	MOTA	16			673	15.609	16.417	21.107	1.00 23.85
20	ATOM	17		PHE		15.213	21.374	20.147	1.00 25.25
	ATOM	18		PHE	673		20.926	19.680	1.00 24.38
	ATOM	19		PHE	673	16.260		19.412	1.00 25.01
	ATOM	20		LEU	674	14.193	21.792		1.00 25.58
	MOTA	21	CA	LEU	674	14.237	21.802	17.969	
25	MOTA	22	CB	LEU	674	12.833	21.974	17.391	1.00 26.05
	ATOM	23	CG	LEU	674	12.067	20.653	17.317	1.00 26.55
	ATOM	24	CD1	LEU	674	10.617	20.887	16.935	1.00 26.35
	ATOM	25	CD2		674	12.762	19.758	16.304	1.00 26.09
	ATOM	26	C	LEU	674	15.199	22.801	17.357	1.00 25.10
30	ATOM	27	ŏ	LEU	674	15.743	22.518	16.294	1.00 26.08
30		28	N	ASN	675	15.440	23.939	18.019	1.00 24.63
	ATOM	29	CA	ASN	675	16.356	24.964	17.484	1.00 23.19
	ATOM			ASN	675	16.478	26.215	18.393	1.00 24.20
	ATOM	30	CB		675	15.206	27.067	18.452	1.00 24.32
25	ATOM	31	CG	ASN		14.368	27.062	17.547	1.00 24.82
35	ATOM	32	OD1		675		27.817	19.539	1.00 24.74
		33	ND2		675	15.076	24.338	17.397	1.00 21.66
	ATOM	34	С	ASN	675	17.726			1.00 21.43
	ATOM	35	0	ASN	675	18.435	24.524	16.417	
	ATOM	36	N	VAL	676	18.095	23.612	18.448	1.00 21.17
40	ATOM	37	CA	VAL	676	19.394	22.952	18.507	1.00 20.92
	ATOM	38	CB	VAL	676	19.718	22.442	19.934	1.00 21.33
	ATOM	39	CG1	VAL	676	18.899	21.237	20.247	1.00 24.09
	ATOM	40	CG2	VAL	676	21.192	22.095	20.065	1.00 21.88
	ATOM	41	Ċ	VAL	676	19.501	21.830	17.473	1.00 19.78
45	ATOM	42	ō	VAL	676	20.421	21.827	16.646	1.00 19.99
40	ATOM	43	N	LEU	677	18.530	20.923	17.434	1.00 19.08
	ATOM	44	CA	LEU	677	18.601	19.848	16.453	1.00 17.91
		45	CB	LEU	677	17.383	18.921	16.518	1.00 17.50
	ATOM				677		18.083	17.798	1.00 16.78
50	MOTA	46		LEU		16.355	16.934	17.541	1.00 17.01
50	ATOM	47		LEU	677		17.555	18.225	1.00 17.10
	ATOM	48		LEU	677	18.615		15.068	1.00 16.96
	ATOM	49	C	LEU	677	18.768	20.427		1.00 14.94
	ATOM	50	0	LEU	677	19.640	20.008	14.347	
	ATOM	51	N	GLU	678	17.980	21.445	14.736	1.00 19.12
55	ATOM	52	CA	GLU	678	18.058	22.121	13.437	1.00 20.06
	ATOM	53	CB	GLU	678	16.972	23.188	13.317	1.00 23.33
	ATOM	54	CG	GLU	678	15.532	22.646	13,381	1.00 28.64
	ATOM	55	CD	GLU	678	14.459	23.736	13.387	1.00 32.31
	ATOM	56		GLU	678	14.811	24.943	13.374	1.00 34.41
60	ATOM	57		GLU	678	13.253	23.384	13.410	1.00 34.91
55	ATOM	58	C	GLU	678	19.410	22.783	13.243	1.00 19.33
	AION	50	Ü	220	5 / 5				

ATON ATON 60 N ALA 679 19.966 23.34 14.329 1.00 18.20 ATON 61 CA ALA 679 21.257 24.018 14.329 1.00 19.45 ATOM 61 CA ALA 679 21.257 24.018 14.329 1.00 19.45 ATOM 63 C ALA 679 21.257 23.094 14.195 1.00 19.45 ATOM 66 C ALA 679 22.472 23.094 14.195 1.00 19.25 ATOM 65 N ILE 680 22.359 21.914 14.802 1.00 18.82 ATOM 66 CA ILE 680 22.359 21.914 14.802 1.00 18.82 ATOM 66 CA ILE 680 23.518 20.994 14.742 1.00 17.49 ATOM 67 CB ILE 680 23.518 20.994 14.742 1.00 17.05 ATOM 69 CGI ILE 680 23.518 19.984 17.159 1.00 15.55 ATOM 70 CDI ILE 680 22.393 19.467 16.391 1.00 15.55 ATOM 71 C ILE 680 23.516 19.984 13.593 1.00 16.89 ATOM 72 O ILE 680 23.516 19.984 13.593 1.00 16.89 ATOM 74 CA GLU 681 22.451 19.992 12.847 1.00 16.89 ATOM 75 CB GLU 681 20.473 16.823 10.348 1.00 16.79 ATOM 76 CG GLU 681 20.473 16.823 10.348 1.00 16.79 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 78 OEL GLU 681 20.599 19.002 11.719 1.00 17.58 ATOM 79 OEL GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 81 O GLU 681 23.370 19.128 10.03 1.30 1.00 17.58 ATOM 82 N PRO 682 23.516 19.984 10.34 10.00 17.83 ATOM 86 C G PRO 682 23.996 16.704 11.00 17.00 17.58 ATOM 87 C OEL GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 20.475 16.502 11.524 1.00 17.58 ATOM 80 C GLU 681 20.475 16.502 11.524 1.00 17.58 ATOM 80 C GLU 681 20.475 16.502 11.524 1.00 17.58 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 80 C GLU 681 23.576 19.128 10.673 1.00 18.79 ATOM 80 C GLU 681 23.576 19.128 10.673 1.00 18.79 ATOM 80 C GLU 681 23.576 19.128 10.673 1.00 18.79 ATOM 100 N VAL 684 26.105 11.524 1.00 17.08 ATOM 100 N VAL 684 26.105 11.524 1.00 17.08 ATOM 90 C GLU 683 25.986 10.00 11.00 10.00 10.00 10.							•			1 00	10 20
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NATION 62							21.257	24.018	14.303		
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AROM	_								14.195	1.00	19.25
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10 ATOM 68 CG2 ILE 680 24.022 21.213 17.159 1.00 16.89 ATOM 70 CD1 ILE 680 22.393 19.467 16.391 1.00 18.55 ATOM 71 C ILE 680 22.558 18.575 17.552 1.00 13.80 ATOM 71 C ILE 680 23.516 19.984 13.593 1.00 16.89 ATOM 72 C ILE 680 24.518 19.303 13.370 1.00 17.12 15 ATOM 73 N GLU 681 22.451 19.922 12.847 1.00 16.79 ATOM 75 CB GLU 681 22.451 19.922 12.847 1.00 16.79 ATOM 75 CB GLU 681 20.579 18.300 9.952 1.00 17.26 ATOM 76 CG GLU 681 20.579 18.300 9.952 1.00 17.58 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 17.58 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 17.58 ATOM 78 OE1 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OE2 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OE2 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OE2 GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.22 ATOM 80 C GEU 681 23.517 20.173 10.043 1.00 20.09 ATOM 80 C GEU 682 25.525 18.021 9.472 1.00 19.12 10.00 19.02 ATOM 80 C GB PRO 682 25.552 18.021 9.472 1.00 19.12 10.00 19.03 ATOM 86 CG PRO 682 25.581 16.546 9.493 1.00 18.30 ATOM 86 CG PRO 682 25.388 16.109 10.846 1.00 17.08 ATOM 80 CG PRO 682 25.388 16.109 10.846 1.00 17.08 ATOM 80 CG PRO 682 25.338 16.109 10.846 1.00 17.00 19.75 ATOM 80 CG PRO 682 25.338 16.109 10.846 1.00 17.00 19.75 ATOM 90 CG LY 683 25.901 18.955 7.339 1.00 20.64 ATOM 90 CG LY 683 25.901 18.955 7.339 1.00 20.64 ATOM 90 CG LY 683 25.901 18.955 7.339 1.00 20.64 ATOM 90 CG LY 683 25.901 18.956 7.339 1.00 20.64 ATOM 90 CG LY 683 25.901 18.956 7.379 1.00 23.13 ATOM 90 CG LY 683 25.901 18.956 7.379 1.00 23.13 ATOM 90 CG LY 684 25.133 19.288 1.131 1.00 23.58 ATOM 91 CG LYAL 685 29.012 14.878 2.665 1.00 17.75 ATOM 90 CG LYAL 685 29.012 14.878 2.665 1.00 17.75 ATOM 90 CG LYAL 685 29.012 14.878 2.665 1				СВ	ILE	680	23.674				
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10					680	24.022	21.213	17.158		
ATOM 70 CDI ILE 680 22.558 18.575 17.552 1.00 13.80 ATOM 71 C ILE 680 23.516 19.984 13.593 1.00 16.89 ATOM 72 O ILE 680 23.516 19.984 13.593 1.00 16.89 ATOM 73 N GLU 681 22.415 19.922 12.847 1.00 16.79 ATOM 74 CA GLU 681 22.455 19.002 11.719 1.00 17.26 ATOM 75 CB GLU 681 20.902 19.227 11.094 1.00 17.26 ATOM 76 CG GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 78 OEL GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OEL GLU 681 20.214 15.981 9.467 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 18.22 ATOM 83 CD PRO 682 23.969 16.704 11.09 1.00 20.09 ATOM 84 CA PRO 682 25.252 18.021 9.472 1.00 19.12 ATOM 85 CB PRO 682 25.381 16.504 10.01 10.10 10.01 18.23 ATOM 86 CG PRO 682 25.385 16.09 10.846 1.00 17.08 ATOM 87 C PRO 682 23.971 18.302 19.472 1.00 19.13 ATOM 88 O PRO 682 23.971 18.302 19.472 1.00 19.13 ATOM 87 C PRO 682 23.971 18.302 19.472 1.00 19.13 ATOM 88 O PRO 682 25.388 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 23.971 18.302 19.472 1.00 19.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 91 C GLY 663 25.595 17.108 5.355 1.00 23.47 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 93 CWAL 684 26.350 18.216 1.259 1.00 24.93 ATOM 94 CA VAL 684 26.350 18.216 1.259 1.00 22.44 ATOM 99 C WAL 684 26.350 18.216 1.259 1.00 22.49 ATOM 99 C WAL 684 26.350 18.216 1.259 1.00 24.93 ATOM 99 C WAL 684 26.350 18.216 1.259 1.00 1.00 23.47 ATOM 99 C WAL 685 29.012 14.978 2.665 1.00 17.86 ATOM 100 C WAL 685 30.303 13.189 4.086 1.00 17.80 ATOM 101 CA WAL 685 30.303 13.189 4.086 1.00 17.76 ATOM 103 CG1 WAL 685 30.030 13.189 4.086 1.00 17.76 ATOM 104 CG2 WAL 685 30.030 13.189 4.086 1.00 17.76 ATOM 105 C WAL 685 30.030 13.189 4.086 1.00 17.76 ATOM 106 C A CYS 686 30.166 1.00 17.55 1.00 17.76 ATOM 107 C A GLA 687 31.161 19.	10						22 393	19.467	16.391	1.00	15.55
ATOM 71 C ILE 680 23.516 19.984 13.593 1.00 16.89 ATOM 72 O ILE 680 24.518 19.303 13.570 1.00 17.12 ATOM 73 N GLU 681 22.415 19.922 12.847 1.00 16.79 ATOM 74 CA GLU 681 22.455 19.002 11.719 1.00 16.79 ATOM 75 CB GLU 681 20.902 19.227 11.094 1.00 16.72 ATOM 76 CG GLU 681 20.957 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 79 OE2 GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 79 OE2 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OE2 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 80 C GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 81 O GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 82 N PRO 682 23.370 19.128 10.673 1.00 18.79 ATOM 83 C GLU 681 20.579 18.300 9.952 1.00 18.79 ATOM 84 CA PRO 682 23.369 16.502 11.524 1.00 17.58 ATOM 85 CB PRO 682 23.969 16.502 11.524 1.00 17.58 ATOM 86 CG PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 86 CG PRO 682 25.525 18.021 9.472 1.00 19.02 ATOM 86 CG PRO 682 25.681 16.546 9.493 1.00 18.30 ATOM 86 CG PRO 682 25.681 16.546 9.493 1.00 18.30 ATOM 87 C PRO 682 23.771 18.392 7.625 1.00 21.67 ATOM 87 C PRO 682 23.771 18.392 7.625 1.00 21.67 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 23.47 ATOM 91 C GLY 683 25.901 18.995 7.339 1.00 23.47 ATOM 92 CO GLY 683 25.901 18.995 7.339 1.00 23.47 ATOM 93 N VAL 684 26.365 17.573 2.685 1.00 21.47 ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 23.47 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 96 CGI VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 97 CG2 VAL 684 25.153 19.228 1.131 1.00 24.89 ATOM 98 C VAL 685 29.912 14.978 2.665 1.00 17.08 ATOM 99 CG VAL 685 29.912 14.978 2.665 1.00 17.05 ATOM 10 CB VAL 685 29.912 14.978 2.665 1.00 17.05 ATOM 10 CG VAL 685 29.912 14.978 2.665 1.00 17.05 ATOM 10 CG VAL 685 29.912 14.978 2.665 1.00 17.05 ATOM 10 CG VAL 685 30.303 13.189 4.086 1.00 17.06 ATOM 10 CG VAL 685 30.303 13.189 4.086 1.00 17.06 ATOM 10 CG VAL 685 30.303 13.189 0.009 1.00 17.00 ATOM 10 CG VAL 685 30.001 11.003 2.259 1.00 16.97 ATOM 10 CG VAL 685 30.001 11.003 2.259 1.0									17.552	1.00	13.80
ATOM 72 O ILE 680 24.518 19.303 13.370 1.00 17.12 ATOM 73 N GLU 681 22.415 19.922 12.847 1.00 16.79 ATOM 74 CA GLU 681 22.415 19.922 12.847 1.00 17.26 ATOM 76 CG GLU 681 20.902 19.227 11.094 1.00 17.26 ATOM 77 CD GLU 681 20.579 18.300 9.952 10.01 8.48 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 78 OEI GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 79 OEI GLU 681 20.579 18.300 9.952 10.0 18.48 ATOM 79 OEI GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OEI GLU 681 20.214 15.981 9.467 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 ATOM 83 CD PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 84 CA PRO 682 25.525 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.11 ATOM 86 CG PRO 682 25.525 18.021 9.472 1.00 19.13 ATOM 87 C PRO 682 23.371 18.382 7.625 1.00 19.08 ATOM 88 O PRO 682 23.371 18.382 7.625 1.00 21.13 ATOM 89 O RA GLY 683 25.901 18.955 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.955 7.339 1.00 20.13 ATOM 90 CA GLY 683 25.901 18.955 7.339 1.00 23.13 ATOM 91 C GLY 683 25.901 18.955 7.339 1.00 23.13 ATOM 92 C GLY 683 25.901 18.260 4.990 1.00 23.13 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.13 ATOM 94 CA VAL 684 26.355 17.573 2.685 1.00 23.47 ATOM 97 CG2 VAL 684 25.153 19.228 1.131 1.00 24.57 ATOM 98 C VAL 684 25.153 19.228 1.131 1.00 24.57 ATOM 99 C CA VAL 684 26.320 18.216 1.259 1.00 24.53 ATOM 99 C VAL 685 29.114 31.189 4.086 1.00 17.88 ATOM 99 C CA GLY 685 30.303 13.189 4.086 1.00 17.88 ATOM 100 C C VAL 685 29.114 31.112 1.3455 1.00 17.76 ATOM 107 N CYS 686 30.451 13.628 -0.650 1.00 17.55 ATOM 108 CA VAL 685 29.114 14.12 1.3455 1.00 17.55 ATOM 109 C C VAL 685 30.031 13.189 4.086 1.00 17.55 ATOM 109 C C VAL 685 30.461 1.167 -0.327 1.00 16.97 ATOM 100 C C VAL 685 30.461 1.167 -0.327 1.00 16.97 ATOM 101 C C VAL 685 30.461 1.167 -0.327 1.00 16.97 ATOM 102 C C VAL 686 30.166 16.031 -2.147 1.00 16.37 ATOM 103 C C VAL 686 30.166 16.		ATOM									
15 ATOM 73 N GLU 681 22.415 19.922 12.847 1.00 16.79 ATOM 74 CA GLU 681 22.265 19.002 11.719 1.00 16.72 ATOM 75 CB GLU 681 20.979 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 78 OE1 GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 80 C GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 81 O GLU 681 23.317 20.173 10.043 1.00 18.79 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 ATOM 83 CD PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 84 CA PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.12 ATOM 86 CG PRO 682 25.538 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 88 O PRO 682 23.91 18.925 7.339 1.00 18.30 ATOM 88 O PRO 682 23.91 18.925 7.339 1.00 18.30 ATOM 89 N GLY 683 25.965 19.422 5.972 1.00 21.13 ATOM 90 CA GLY 683 25.965 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.965 17.108 5.335 1.00 23.47 ATOM 92 O GLY 683 25.965 17.108 5.335 1.00 23.47 ATOM 93 N VAL 684 26.320 18.216 1.259 1.00 21.47 ATOM 96 CGI VAL 684 26.320 18.216 1.259 1.00 23.47 ATOM 97 CC2 VAL 684 26.320 18.216 1.259 1.00 23.47 ATOM 98 C VAL 684 26.320 18.216 1.259 1.00 23.47 ATOM 98 C VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 99 C CA GLY 683 25.955 17.108 5.335 1.00 23.47 ATOM 99 C CA GLY 683 25.955 17.108 2.311 1.00 24.89 40 ATOM 98 C VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 99 C CA GLY 685 28.955 17.108 2.665 1.00 17.80 ATOM 99 C CA GLY 685 28.955 17.108 1.00 18.50 ATOM 100 N VAL 685 29.915 13.8957 2.665 1.00 17.80 ATOM 107 N CYS 686 30.124 14.339 0.609 1.00 17.55 ATOM 108 CA CYS 686 30.124 14.339 0.609 1.00 17.55 ATOM 109 CB CYS 686 30.141 12.496 0.615 1.00 17.74 ATOM 110 CA VAL 685 29.915 13.8577 1.055 1.00 17.75 ATOM 110 CA VAL 685 30.303 1.161 8.929 -1.295 1.00 16.91 ATOM 110 CA VAL 685 30.303 1.161 8.929 -1.295 1.00 16.91 ATOM 110 CA VAL 685 30.303 1.161 8.929 -1.295 1.00 16.91 ATOM 110 CA VAL 685 30.303 1.161 8.929 -1.295 1.00 16.91 ATOM 1		ATOM	71	С							
ATOM 75 CB GLU 681 22.265 19.002 11.719 1.00 17.26 ATOM 75 CB GLU 681 20.902 19.227 11.094 1.00 16.72 ATOM 76 CG GLU 681 20.579 18.300 9.952 1.00 18.48 1.00 17.51 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 1.00 17.51 ATOM 78 OE1 GLU 681 20.679 16.502 11.524 1.00 17.51 ATOM 79 OE2 GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 80 C GLU 681 20.214 15.981 9.467 1.00 18.79 ATOM 81 O GLU 681 23.517 20.173 10.043 1.00 20.09 ATOM 82 N PRO 682 23.370 19.128 10.6673 1.00 18.79 ATOM 84 CD PRO 682 23.999 16.704 11.019 1.00 18.22 ATOM 84 CD PRO 682 23.999 16.704 11.019 1.00 18.22 ATOM 85 CB PRO 682 25.581 16.546 9.493 1.00 18.30 ATOM 86 CG PRO 682 25.583 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 23.571 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 23.47 ATOM 91 C GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 95 CB VAL 684 26.300 18.266 4.990 1.00 23.47 ATOM 97 CG2 VAL 684 26.300 18.266 1.259 1.00 24.54 ATOM 97 CG2 VAL 684 26.300 18.266 1.259 1.00 24.54 ATOM 97 CG2 VAL 684 26.300 18.266 1.259 1.00 24.57 ATOM 98 CVAL 684 26.300 18.266 1.259 1.00 24.57 ATOM 97 CG2 VAL 684 26.300 18.266 1.259 1.00 24.57 ATOM 98 CVAL 684 26.301 18.216 1.259 1.00 24.57 ATOM 97 CG2 VAL 684 26.301 18.216 1.259 1.00 24.57 ATOM 97 CG2 VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 98 C VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 97 CG2 VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 97 CG2 VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.81 ATOM 100 CA VAL 685 29.012 14.978 2.665 1.00 17.85 ATOM 100 CA VAL 685 29.013 14.112 1.345 1.00 17.55 ATOM 110 CA VAL 685 30.303 13.189 1.00 6.99 1.00 17.85 ATOM		ATOM	72	0	ILE						
ATOM 74 CA GLU 681 22.265 19.002 11.79 1.00 16.72 ATOM 75 CB GLU 681 20.902 19.227 11.094 1.00 16.72 ATOM 76 CG GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 78 0E1 GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 79 0E2 GLU 681 20.473 16.823 10.348 1.00 17.58 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 82 N PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 83 CD PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.12 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.12 ATOM 86 CG PRO 682 25.538 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 23.711 18.382 7.625 1.00 19.76 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 21.13 ATOM 90 CA GLY 683 25.595 17.108 5.355 1.00 21.13 ATOM 91 C GLY 683 25.595 17.108 5.355 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.13 ATOM 94 CA VAL 684 26.320 18.216 1.259 1.00 24.44 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 96 CG1 VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 99 O VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 99 C CVAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 102 CB VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 103 CG1 VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 104 CG2 VAL 685 28.238 13.367 0.969 1.00 17.81 ATOM 105 C VAL 685 28.238 13.367 0.969 1.00 17.55 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.57 ATOM 108 CA CYS 686 30.3151 12.496 0.615 1.00 17.57 ATOM 110 CC VS 686 30.3151 1.324 2.965 1.00 17.55 ATOM 110 CC VS 686 30.3151 1.324 2.965 1.00 17.55 ATOM 110 CA ALA 687 31.161 13.599 -1.295 1.00 16.97 ATOM 110 CC S ALA 687 31.161 13.699 -1.295 1.00 16.97 ATOM 110 CC S ALA 687 31.161 13.699 -1.310 1.00 17.55 ATOM 111 CC CS 686 31.3161 12.496 0.615 1.00 17.56 ATOM 111 CC CS 666 31.3166 1.00 17.59 1.00 17.59	15	ATOM	73	N	GLU	681					
ATOM 76 CB GLU 681 20.902 19.227 11.094 1.00 16.72 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.51 17.51 ATOM 79 OE2 GLU 681 20.214 15.981 9.467 1.00 18.59 ATOM 80 C GLU 681 20.214 15.981 9.467 1.00 18.59 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 20.09 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 20.09 ATOM 84 CA PRO 682 25.522 18.021 9.472 1.00 19.11 1.00 18.22 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.11 1.00 18.22 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.11 1.00 18.30 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 19.17 0.00 19.70 1.00 18.30 ATOM 88 O PRO 682 25.338 16.109 10.846 1.00 19.76	. •		74	CA	GLU	681	22.265	19.002			
ATOM 76 CG GLU 681 20.579 18.300 9.952 1.00 18.48 ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 78 0E1 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 0E2 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 ATOM 85 CB PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 85 CB PRO 682 25.525 18.021 9.472 1.00 19.12 ATOM 86 CG PRO 682 25.525 18.021 9.472 1.00 19.12 ATOM 86 CG PRO 682 25.538 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 25.538 16.109 10.846 1.00 17.08 ATOM 88 O PRO 682 23.979 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 91 C GLY 683 25.595 17.108 5.355 1.00 21.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 94 CA VAL 684 26.309 18.260 4.990 1.00 23.13 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.44 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 99 C VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 99 C VAL 684 26.320 18.216 1.259 1.00 24.45 ATOM 99 C VAL 684 25.153 19.228 1.131 1.00 24.45 ATOM 99 C VAL 684 25.153 19.228 1.131 1.00 24.45 ATOM 99 C VAL 685 27.778 15.631 2.5655 1.00 17.81 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 102 CB VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 103 CG1 VAL 685 29.012 14.878 2.665 1.00 17.85 ATOM 105 C VAL 685 29.012 14.878 2.665 1.00 17.05 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.05 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.05 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.05 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.05 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.55 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.55 ATOM 107 CA VAL 685 29.012 14.878 2.665 1.00 17.00 17.55 ATOM 107 CA VAL 685 28.238 13.367 0.060 1.00 17.55 ATOM 107 CA VAL 685 33.370 1							20.902	19.227	11.094		
ATOM 77 CD GLU 681 20.473 16.823 10.348 1.00 17.51 ATOM 78 OEI GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OE2 GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.59 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 10.40								18.300	9.952	1.00	18.48
ATOM 78 OEI GLU 681 20.659 16.502 11.524 1.00 17.58 ATOM 79 OE2 GLU 681 20.214 15.981 9.467 1.00 18.59 ATOM 80 C GLU 681 23.517 20.173 10.043 1.00 20.09 ATOM 81 O GLU 681 23.517 20.173 10.043 1.00 20.09 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 ATOM 85 CB PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 85 CB PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 86 CG PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.11 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.16 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.16 ATOM 87 C PRO 682 25.681 16.546 9.493 1.00 19.70 19										1.00	17.51
ATOM 79 OE2 GLU 681 20.214 15.981 9.467 1.00 18.59 ATOM 80 C GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 81 O GLU 681 23.370 19.128 10.673 1.00 18.79 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 1.00 19.02 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.02 1.00 18.22 ATOM 84 CA PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.681 16.546 9.493 1.00 18.30 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 89 N GLY 683 25.665 19.422 5.972 1.00 19.76 ATOM 90 CA GLY 683 25.809 18.260 4.990 10.00 23.13 ATOM 91 C GLY 683 25.809 18.260 4.990 10.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.54 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 22.44 ATOM 95 CB VAL 684 26.365 17.573 2.665 1.00 22.44 ATOM 96 CG1 VAL 684 26.365 17.573 2.665 1.00 22.44 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 99 O VAL 684 26.350 18.216 1.259 1.00 24.93 ATOM 99 O VAL 684 25.539 17.108 5.355 1.00 24.93 ATOM 99 CG2 VAL 684 25.539 17.109 2.865 1.00 24.89 ATOM 97 CG2 VAL 684 25.539 17.103 0.183 1.00 24.89 ATOM 101 CA VAL 685 27.778 15.631 2.585 1.00 19.82 ATOM 102 CB VAL 685 27.778 15.631 2.585 1.00 19.82 ATOM 103 CG1 VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 103 CG1 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 105 C VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 107 CC3 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 108 CA CYS 686 30.451 13.662 -0.650 1.00 17.55 ATOM 107 CC3 VAL 685 28.955 13.857 3.867 1.00 17.55 ATOM 107 CC3 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 107 CC3 CAL 685 28.955 13.857 3.867 1.00 17.55 ATOM 107 CC3 CAL 686 30.451 13.662 -0.650 1.00 17.55 ATOM 107 CC3 CAL 686 30.451 13.662 -0.650 1.00 17.55 ATOM 107 CC3 CAL 686 30.166 16.031 -2.147 -0.0327 1.00 16.97 ATOM 107 CC3 CAL 686 30.166 16.031 -2.147 -0.0327 1.00 16.97 ATOM 107 CC3 CAL 686 30.166 16.031 -2.147 -0.0327 1.00 16.97 ATOM 107 CC3 CAL 686 30.166 16.031 -2.147 -0.0327 1.00 16.97 ATOM											
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ATOM 81 0 GLU 681 23.517 20.173 10.043 1.00 20.09 ATOM 82 N PRO 682 24.145 18.044 10.437 1.00 19.22 ATOM 83 CD PRO 682 23.969 16.704 11.019 1.00 19.22 ATOM 84 CA PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 24.912 18.475 8.057 1.00 19.76 ATOM 87 C PRO 682 23.771 18.382 7.625 1.00 21.67 ATOM 99 N GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 90 CA GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 92 O GLY 683 25.655 19.422 5.972 1.00 21.67 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 95 CB VAL 684 26.365 17.573 2.685 1.00 22.44 ATOM 96 CG1 VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 98 C VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 98 C VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 99 O VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 99 O VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 99 O VAL 684 26.216 19.034 1.311 1.00 24.89 ATOM 99 O VAL 684 28.708 17.614 3.042 1.00 19.82 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 100 N VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 103 CG1 VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 103 CG1 VAL 685 29.12 14.314 1.112 1.345 1.00 15.58 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 105 C VAL 685 28.257 14.556 5.147 1.00 15.58 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 17.70 17.89 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.75 1.55 1.00 17.76 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.76 ATOM 109 CB CYS 686 30.146 16.031 -2.147 1.00 21.38 ATOM 107 N CYS 686 30.146 16.031 -2.147 1.00 11.76 1.76 ATOM 110 SG CYS 686 31.354 12.447 -0.327 1.00 16.91 ATOM 110 SG CYS 686 31.354 12.447 -0.327 1.00 16.91 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.76 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.76 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.76 ATOM		ATOM	79	OE2				_			
ATOM 81 0 GLU 681 23.517 20.173 10.0437 1.00 19.09 ATOM 82 N PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 83 CD PRO 682 23.969 16.704 11.019 1.00 18.22 ATOM 84 CA PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 88 O PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 91 C GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.190 18.567 3.748 1.00 23.48 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 96 CGI VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 26.217 17.130 0.183 1.00 24.59 ATOM 98 C VAL 684 26.217 17.130 0.183 1.00 24.59 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 24.89 ATOM 99 C VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 C VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 C VAL 685 27.778 15.631 2.28 1.131 1.00 24.89 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 105 C VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 106 O VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 107 N CYS 686 30.24 14.339 0.609 1.00 17.89 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.76 ATOM 108 CYS 686 30.451 13.628 -0.650 1.00 17.76 ATOM 108 CG VS 686 30.451 13.628 -0.650 1.00 17.76 ATOM 108 CG CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.76 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.76 ATOM 115 CB ALA 687 31.183 11.360 -1.055 1.00 17.74 ATOM 116 C ALA 687 31.183 11.360 -1.055 1.00 17.16		ATOM	80	С	GLU	681					
25 ATOM 83 CD PRO 682 23.969 16.704 11.019 1.00 19.22 ATOM 84 CA PRO 682 23.969 16.704 11.019 1.00 19.22 ATOM 85 CB PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 24.912 18.475 8.057 1.00 19.76 ATOM 88 O PRO 682 23.771 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.95 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.955 17.108 5.355 1.00 23.47 ATOM 92 O GLY 683 25.955 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.665 1.00 24.93 ATOM 95 CB VAL 684 26.365 17.573 2.665 1.00 24.93 ATOM 96 CGI VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 97 CG2 VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 98 C VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 99 O VAL 684 26.190 18.567 3.748 1.00 24.59 ATOM 99 O VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 99 O VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 24.64 ATOM 99 O VAL 684 27.778 15.631 2.585 1.00 19.62 ATOM 100 N VAL 685 29.102 14.576 5.147 1.00 17.89 ATOM 101 CA VAL 685 29.102 14.576 5.147 1.00 17.89 ATOM 103 CGI VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 15.27 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 15.58 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.80 ATOM 108 CG CYS 686 30.451 13.628 -0.650 1.00 17.76 ATOM 109 CB CYS 686 30.156 16.031 -2.147 1.00 16.97 ATOM 110 SG CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 111 C C ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 112 C CB ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 116 C ALA 687 31.183 11.360 -1.055 1.00 17.76 ATOM 117 O ALA 687 31.185 9.431 -1.139 1.00 10.91 60 ATOM 118 N GLY 688 33.370 11.1023 -2.559 1.00 18.50			81	0	GLU	681	23.517	20.173			
25 ATOM 83 CD PRO 682 23,969 16.704 11.019 1.00 18.22 ATOM 84 CA PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.681 16.546 9.493 1.00 19.11 0.00 18.30 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 24.912 18.475 8.057 1.00 19.76 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.13 ATOM 93 N VAL 684 26.365 17.573 2.655 1.00 23.47 ATOM 95 CB VAL 684 26.365 17.573 2.655 1.00 23.47 ATOM 95 CB VAL 684 26.365 17.573 2.655 1.00 22.44 ATOM 95 CB VAL 684 26.365 17.573 2.655 1.00 24.57 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 98 C VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 99 C VAL 684 26.320 18.216 1.259 1.00 24.57 ATOM 99 C VAL 684 25.153 19.228 1.131 1.00 24.57 ATOM 99 C VAL 684 25.153 19.228 1.131 1.00 24.57 ATOM 99 C VAL 684 25.755 17.614 3.042 1.00 19.82 ATOM 99 C VAL 685 27.725 16.934 2.811 1.00 24.57 ATOM 101 CA VAL 685 27.778 15.631 2.585 1.00 19.05 ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 105 C VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.58 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.52 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.55 ATOM 108 CA CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 110 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 115 CB ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 116 C ALA 688 33.370 11.023 -2.539 1.00 18.50 ATOM 116 C ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 116 C ALA 688 33.370 11.023 -2.539 1.00 18.50 ATOM 116 C ALA 687 31.185 9.431 -1.139 1.00 17.98 ATOM 117 O ALA 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 68							24.145	18.044	10.437		
ATOM 84 CA PRO 682 25.252 18.021 9.472 1.00 19.11 ATOM 85 CB PRO 682 25.681 16.546 9.493 1.00 18.30 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 88 O PRO 682 23.771 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 94 CA VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 95 CB VAL 684 26.365 17.573 2.685 1.00 24.93 ATOM 96 CG1 VAL 684 26.365 17.573 2.685 1.00 24.93 ATOM 97 CG2 VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 98 C VAL 684 25.153 19.228 1.131 1.00 24.89 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 685 27.778 15.631 2.585 1.00 19.85 ATOM 100 N VAL 685 27.778 15.631 2.585 1.00 19.85 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 28.955 13.857 3.857 1.00 17.89 ATOM 103 CG1 VAL 685 28.955 13.857 3.857 1.00 17.89 ATOM 104 CG2 VAL 685 28.955 13.857 3.857 1.00 17.89 ATOM 105 C VAL 685 28.252 14.556 5.107 17.00 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.81 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.85 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.85 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.85 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.76 ATOM 110 CA ALA 687 31.183 11.360 -1.065 1.00 17.76 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 115 CB ALA 687 31.185 9.431 -1.139 1.00 17.76 ATOM 116 C ALA 687 31.185 9.431 -1.139 1.00 17.76 ATOM 117 O ALA 687 31.185 9.431 -1.139 1.00 17.79 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50	25						23,969	16.704	11.019	1.00	18.22
ATOM 85 CB PRO 682 25.681 16.546 9.493 1.00 18.30 ATOM 86 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM 87 C PRO 682 24.912 18.475 8.057 1.00 19.76 ATOM 88 O PRO 682 23.771 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 99 CA GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 23.47 ATOM 95 CB VAL 684 26.365 17.573 2.685 1.00 23.47 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 98 C VAL 684 27.725 16.934 2.811 1.00 24.89 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 24.89 ATOM 100 N VAL 685 27.778 15.631 2.885 1.00 19.05 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.88 ATOM 108 CA CYS 686 30.451 31.628 -0.650 1.00 17.88 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.86 ATOM 109 CB CYS 686 30.451 31.628 -0.650 1.00 17.55 ATOM 109 CB CYS 686 30.451 13.628 -0.650 1.00 17.75 ATOM 110 CA VAL 687 31.183 11.360 -1.065 1.00 17.74 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 115 CB ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 116 C ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 117 O ALA 687 31.185 9.431 -1.139 1.00 16.97 ATOM 116 C ALA 687 31.185 9.431 -1.139 1.00 16.97 ATOM 117 O ALA 687 31.185 9.431 -1.139 1.00 18.03 ATOM 116 C ALA 687 31.185 9.431 -1.139 1.00 18.06 ATOM 117 O ALA 688 33.370 11.023 -2.539 1.00 18.06	25								9.472	1.00	19.11
ATOM 85 CG PRO 682 25.338 16.109 10.846 1.00 17.08 ATOM ATOM 87 C PRO 682 24.912 18.475 8.057 1.00 19.76 ATOM 88 O PRO 682 23.771 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 22.44 ATOM 95 CB VAL 684 26.365 17.573 2.685 1.00 22.44 ATOM 96 CGI VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 98 C VAL 684 26.217 17.130 0.183 1.00 24.89 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 28.708 17.614 3.042 1.00 19.82 ATOM 100 CA VAL 685 29.012 14.878 2.665 1.00 19.05 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.88 ATOM 102 CB VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 103 CG1 VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 105 C VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 106 O VAL 685 28.955 13.857 3.867 1.00 17.86 ATOM 107 N CYS 686 30.224 14.339 0.699 1.00 18.33 ATOM 107 N CYS 686 30.224 14.339 0.699 1.00 17.08 ATOM 109 CB CYS 686 30.241 13.628 -0.650 1.00 17.55 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 110 CCYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 110 CCYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 111 CC CYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 112 C CYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 115 CB ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 115 CB ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 115 CB ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 115 CB ALA 687 31.185 9.431 -1.139 1.00 17.55 ATOM 115 CB ALA 687 33.377 10.161 -1.526 1.00 18.06 ATOM 115 CB ALA 687 33.377 10.161 -1.526 1.00 18.50 ATOM 115 CB ALA 687 33.377 10.161 -1.526 1.00 18.50 ATOM 117 O ALA 687 33.377 10.161 -1.526 1.00 18.50 ATOM 119 CB ALA 687 33.370 11.002 -2.539 1.00 18.50 ATOM 119 CB ALA 687											
ATOM 86 CG PRO 682 24.912 18.475 8.057 1.00 19.76 ATOM 88 O PRO 682 23.771 18.382 7.625 1.00 21.13 ATOM 89 N GLY 683 25.901 18.995 7.339 1.00 20.64 ATOM 90 CA GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.665 1.00 22.44 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 96 CG1 VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 26.321 17.130 0.183 1.00 24.57 ATOM 98 C VAL 684 25.153 19.228 1.131 1.00 24.89 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 685 27.778 15.631 2.585 1.00 19.82 ATOM 100 N VAL 685 27.778 15.631 2.585 1.00 17.89 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 103 CG1 VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 104 CG2 VAL 685 29.12 14.878 2.665 1.00 17.80 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.80 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 17.80 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.80 ATOM 108 CA CYS 686 30.224 14.339 0.609 1.00 17.00 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.55 ATOM 109 CB CYS 686 30.166 16.031 -2.147 1.00 16.97 ATOM 110 CA VS 686 32.141 12.496 0.615 1.00 17.76 ATOM 111 C CYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 112 O CYS 686 31.354 12.497 -0.327 1.00 16.97 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50		ATOM									
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ATOM 90 CA GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.665 19.422 5.972 1.00 21.67 ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 35 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 22.44 ATOM 95 CB VAL 684 26.365 17.573 2.685 1.00 24.493 ATOM 96 CG1 VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 25.153 19.228 1.131 1.00 24.57 ATOM 98 C VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 685 27.778 15.631 2.585 1.00 19.85 ATOM 100 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.81 ATOM 103 CG1 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 28.257 14.556 5.147 1.00 16.27 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.80 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.50 ATOM 109 CB CYS 686 30.224 14.339 0.609 1.00 17.50 ATOM 109 CB CYS 686 30.166 16.031 -2.147 1.00 16.27 ATOM 109 CB CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 109 CB CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 112 O CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 117 O ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.56	30	ATOM	88	0	PRO	682					
ATOM 91 C GLY 683 25.665 19.422 5.972 1.00 21.6/ ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 35 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 22.44 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 96 CG1 VAL 684 26.310 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 25.153 19.228 1.131 1.00 24.89 40 ATOM 98 C VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 28.708 17.614 3.042 1.00 19.82 ATOM 100 N VAL 685 27.778 15.631 2.585 1.00 19.05 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 19.05 ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 104 CG2 VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 16.27 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 16.27 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 28.913 13.67 0.969 1.00 16.27 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.80 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.82 ATOM 109 CB CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 110 SG CYS 686 31.101 14.534 -1.706 1.00 17.70 ATOM 110 CA ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 111 C CYS 686 32.141 12.496 0.615 1.00 17.75 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.75 ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 115 CB ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 116 C ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 116 C ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 116 C ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 116 C ALA 687 31.949 10.132 -0.836 1.00 17.75 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 16.91 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 33.370 11.023 -2.539 1.00 18.50	-			N	GLY	683	25.901	18.995	7.339		
ATOM 91 C GLY 683 25.809 18.260 4.990 1.00 23.13 ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 22.44 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 97 CG2 VAL 684 26.217 17.130 0.183 1.00 24.89 40 ATOM 98 C VAL 684 25.153 19.228 1.131 1.00 20.64 ATOM 99 O VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 28.708 17.614 3.042 1.00 19.82 ATOM 100 N VAL 685 27.778 15.631 2.585 1.00 19.82 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.80 ATOM 108 CA CYS 686 30.224 14.339 0.609 1.00 17.80 ATOM 109 CB CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.52 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 110 CA VAL 687 31.183 11.360 -1.065 1.00 17.55 ATOM 111 C CYS 686 32.141 12.496 0.615 1.00 17.55 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.55 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 17.55 ATOM 115 CB ALA 687 33.277 10.161 -1.526 1.00 17.55 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 17.57 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 17.55 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50						683	25.665	19.422	5.972		
ATOM 92 O GLY 683 25.595 17.108 5.355 1.00 23.47 ATOM 93 N VAL 684 26.190 18.567 3.748 1.00 23.58 ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 24.49 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 96 CG1 VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 97 CG2 VAL 684 25.153 19.228 1.131 1.00 24.89 40 ATOM 98 C VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 28.708 17.614 3.042 1.00 19.82 ATOM 100 N VAL 685 27.778 15.631 2.585 1.00 19.05 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 103 CG1 VAL 685 28.955 13.857 3.867 1.00 15.58 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.88 ATOM 108 CA CYS 686 30.224 14.339 0.609 1.00 17.05 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.55 ATOM 110 CCYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.55 ATOM 115 CB ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 116 C ALA 687 31.183 11.360 -1.065 1.00 17.75 ATOM 117 O ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50							25.809	18.260	4.990	1.00	23.13
ATOM										1.00	23.47
ATOM 94 CA VAL 684 26.365 17.573 2.685 1.00 22.44 ATOM 95 CB VAL 684 26.320 18.216 1.259 1.00 24.93 ATOM 96 CG1 VAL 684 26.217 17.130 0.183 1.00 24.57 ATOM 97 CG2 VAL 684 25.153 19.228 1.131 1.00 24.89 40 ATOM 98 C VAL 684 27.725 16.934 2.811 1.00 20.64 ATOM 99 O VAL 684 28.708 17.614 3.042 1.00 19.82 ATOM 100 N VAL 685 27.778 15.631 2.585 1.00 19.05 ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.89 ATOM 103 CG1 VAL 685 30.303 13.189 4.086 1.00 17.58 ATOM 104 CG2 VAL 685 30.303 13.189 4.086 1.00 15.58 ATOM 105 C VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 106 O VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.88 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.74 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 115 CB ALA 687 31.181 11.360 -1.055 1.00 17.74 ATOM 116 C ALA 687 31.183 11.360 -1.055 1.00 17.75 ATOM 116 C ALA 687 31.183 11.360 -1.055 1.00 17.74 ATOM 115 CB ALA 687 31.183 11.360 -1.055 1.00 17.75 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50	`^=										
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ATOM 101 CA VAL 685 29.012 14.878 2.665 1.00 17.89 ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.81 45 ATOM 103 CG1 VAL 685 30.303 13.189 4.086 1.00 15.58 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 28.238 13.367 0.969 1.00 17.88 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.00 50 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.74 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 33.370 11.023 -2.539 1.00 18.50 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16									2.585	1.00	19.05
ATOM 102 CB VAL 685 28.955 13.857 3.867 1.00 17.81 ATOM 103 CG1 VAL 685 30.303 13.189 4.086 1.00 15.58 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 28.238 13.367 0.969 1.00 18.33 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.00 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.75 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16										1.00	17.89
45 ATOM 103 CG1 VAL 685 30.303 13.189 4.086 1.00 15.58 ATOM 104 CG2 VAL 685 28.527 14.556 5.147 1.00 16.27 ATOM 105 C VAL 685 29.143 14.112 1.345 1.00 17.88 ATOM 106 O VAL 685 28.238 13.367 0.969 1.00 18.33 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.00 17.00 ALA 685 ATOM 110 SG CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.15 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 115 CB ALA 687 33.277 10.161 -1.526 1.00 17.98 ATOM 117 O ALA 687 33.277 10.161 -1.526 1.00 17.98 ATOM 117 O ALA 687 33.370 11.023 -2.539 1.00 18.06 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16											
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ATOM 106 O VAL 685 28.238 13.367 0.969 1.00 18.33 ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.00 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 16.97 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.74 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16			105	С	VAL	685	29.143				
ATOM 107 N CYS 686 30.224 14.339 0.609 1.00 17.00 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.15 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16						685	28.238	13.367	0.969	1.00	18.33
50 ATOM 108 CA CYS 686 30.451 13.628 -0.650 1.00 17.52 ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.15 55 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 118 N <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td>30.224</td><td>14.339</td><td>0.609</td><td>1.00</td><td>17.00</td></td<>							30.224	14.339	0.609	1.00	17.00
ATOM 109 CB CYS 686 31.101 14.534 -1.706 1.00 17.76 ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.15 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16	50									1.00	17.52
ATOM 110 SG CYS 686 30.166 16.031 -2.147 1.00 21.38 ATOM 111 C CYS 686 31.354 12.447 -0.327 1.00 16.97 ATOM 112 O CYS 686 32.141 12.496 0.615 1.00 17.15 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16	50										
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55 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16		ATOM	111	С	CYS					1.00	10.57
55 ATOM 113 N ALA 687 31.183 11.360 -1.065 1.00 17.74 ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16		ATOM	112	0	CYS	686					
ATOM 114 CA ALA 687 31.949 10.132 -0.836 1.00 17.57 ATOM 115 CB ALA 687 31.161 8.929 -1.295 1.00 16.91 ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16	55			N	ALA	687	31.183	11.360			
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ATOM 116 C ALA 687 33.277 10.161 -1.526 1.00 18.06 ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16										1.00	16.91
ATOM 117 O ALA 687 34.185 9.431 -1.139 1.00 17.98 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16											
60 ATOM 118 N GLY 688 33.370 11.023 -2.539 1.00 18.50 ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16											
ATOM 119 CA GLY 688 34.580 11.167 -3.326 1.00 19.16											
A10H 115 CA CD: 300 1000	60										
		ATOM	119	CA		688					
				С	GLY	688	34.705	10.099	-4.388	1.00	19.90

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	ATOM ATOM	121 122	0 N	GLY HIS	688 689	35.802 33.582	9.630	-4.771 -4.907	1.00 2 1.00 2 1.00 2	20.92
	ATOM	123	CA	HIS	689	33.57		-5.912	1.00 2	22.33
	ATOM	124	CB	HIS	689	32.19		-5.900 -6.857	1.00 2	
5	ATOM	125	CG	HIS	689	32.04		-7.033	1.00	22 64
•	ATOM	126	CD2		689	32.78		-7.796	1.00	
	ATOM	127	ND1		689	31.04		-8.516	1.00	
	ATOM	128	CEl		689	31.16		-8.074	1.00	
	ATOM	129	NE2	HIS	689	32.21		-7.328	1.00	
10	ATOM	130	С	HIS	689	33.92		-7.731	1.00	
	ATOM	131	0	HIS	689	33.51		-8.073	1.00	26.27
	ATOM	132	N	ASP	690	34.71		-9.447	1.00	28.86
	ATOM	133	CA	ASP	690	35.01		-9.963	1.00	
	ATOM	134	CB	ASP	690	36.33		-11.361	1.00	30.03
15	MOTA	135	CG	ASP	690	36.69		-11.764	1.00	31.23
	MOTA	136		ASP	690	37.86		-12.061	1.00	
	ATOM	137		ASP	690	35.81 33.87		-10.286	1.00	
	MOTA	138	С	ASP	690	33.70			1.00	
	MOTA	139	0	ASP	690	33.76		-10.832	1.00	
20	MOTA	140	N	ASN	691	31.93				33.60
	ATOM	141	CA	ASN	691 691	30.72		-11.416	1.00	
	MOTA	142	CB	ASN	691	30.07	-		1.00	
	ATOM	143	CG	ASN ASN	691	29.18	• • • • • •	-9.930	1.00	32.13
0.5	ATOM	144		ASN	691	30.54			1.00	33.57
25	ATOM	145		ASN	691	32.28		-13.136		35.13
	ATOM	146	C O	ASN	691	31.41	9 8.733	-13.999		36.66
	ATOM	147 148	N	ALA	692	33.56	5 8.471	-13.434		36.15
	ATOM	149	CA	ALA	692	33.99	8.365	-14.819		37.44
30	ATOM ATOM	150	CB	ALA	692	35.14	8 9.328	-15.103		36.71
30	ATOM	151	Č	ALA	692	34.42		-15.027		38.39
	ATOM	152	ō	ALA	692	34.41		-16.139		39.10
	ATOM	153	N	GLN	693	34.75		-13.928		39.29
	ATOM	154	CA	GLN	693	35.20		-13.942	1.00	40.00
35	ATOM	155	CB	GLN	693	36.13		-12.745		41.81
	MOTA	156	CG	GLN	693	37.53		-13.110		44.34
	ATOM	157	CD	GLN	693	38.42		-13.902		45.44 45.95
	ATOM	158	OE1		693	39.3		-13.363		45.47
	ATOM	159	NE2	GLN	693	38.1		3 -15.186		39.48
40	ATOM	160	С	GLN	693	33.98		9 -13.854 3 -13.217		40.11
	ATOM	161	0	GLN	693	32.99	_	3 -13.217		38.78
	ATOM	162	Ŋ	PRO	694	34.0		5 -15.375		38.88
	ATOM	163	CD	PRO	694	35.13		2 -14.489		36.98
	ATOM	164	CA	PRO	694	32.9° 33.5°		17.365		37.17
45	ATOM	165	CB	PRO	694	34.4		1 -16.234		38.48
	ATOM	166	CG	PRO	694 694	32.5		-13.109		35.56
	MOTA	167	C	PRO PRO	694	33.4		3 -12.204	1.00	35.44
	ATOM	168	0	ASP	695	31.2		2 -12.958	1.00	34.27
50	ATOM	169	N CA	ASP	695	30.7		4 -11.698		32.38
50	ATOM	170 171	CB	ASP	695	29.2		8 -11.694		29.77
	ATOM	172	CG	ASP	695	28.6		1 -11.608	1.00	28.80
	MOTA MOTA	173		1 ASP	695	27.5		0 -12.089		27.09
	ATOM	174		2 ASP	695	29.3	29 2.79	4 -11.057		28.72
55	ATOM	175	C.	ASP	695	31.3	18 -0.86	8 -11.524		32.55
55	ATOM	176	ō	ASP	695	31.2		7 -12.429		33.50
	ATOM	177	N	SER	696	32.0		2 -10.424		31.71
	ATOM	178	CA		696	32.5	77 -2.33	3 -10.077		30.42
	ATOM	179	СВ		696	34.0	64 -2.38	3 -10.425		30.43
60	ATOM	180	ŌĞ		696	34.8		9 -9.567		31.47
	ATOM	181	C	SER	696	32.3	40 -2.44			30.15
	ATOM	182	0	SER	696	32.2	75 -1.41	8 -7.885	1.00	30.10

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ATOM 183 N PHE 697 31.890 -3.933 -6.679 1.00 26.74 ATOM 186 CG PHE 697 31.982 -5.442 -6.623 1.00 25.74 ATOM 186 CG PHE 697 31.982 -5.442 -4.989 1.00 24.31 ATOM 187 CD1 PHE 697 30.536 -5.722 -4.988 1.00 24.31 ATOM 188 CD2 PHE 697 30.545 -6.281 -4.220 1.00 24.78 ATOM 199 CE2 PHE 697 30.344 -6.063 -3.071 1.00 24.68 ATOM 199 CE2 PHE 697 30.344 -6.063 -3.071 1.00 24.67 ATOM 191 CZ PHE 697 30.344 -6.063 -3.071 1.00 24.67 ATOM 191 CZ PHE 697 31.406 -6.512 -2.315 1.00 24.37 ATOM 192 C PHE 697 31.406 -6.512 -2.315 1.00 24.37 ATOM 193 O PHE 697 32.641 -2.393 -4.972 1.00 24.37 ATOM 194 N ALA 698 34.219 -3.495 -6.140 1.00 25.76 ATOM 195 CA ALA 698 35.351 -2.911 -5.436 1.00 25.76 ATOM 197 C ALA 698 35.351 -2.911 -5.436 1.00 25.52 ATOM 199 N ALA 698 35.351 -2.911 -5.436 1.00 25.52 ATOM 199 N ALA 698 35.323 -1.402 -5.300 1.00 25.52 ATOM 199 N ALA 698 35.323 -1.402 -5.300 1.00 25.52 ATOM 199 N ALA 699 35.029 -0.737 -6.414 1.00 25.51 ATOM 199 N ALA 699 34.845 1.156 -7.943 1.00 25.51 ATOM 201 CB ALA 699 34.845 1.156 -7.943 1.00 25.51 ATOM 202 C ALA 699 34.845 1.156 -7.943 1.00 25.51 ATOM 203 C A ALA 699 34.845 1.156 -7.943 1.00 24.31 ATOM 204 N ALA 699 34.845 1.156 -7.943 1.00 24.31 ATOM 205 CB ALA 699 34.845 1.156 -7.943 1.00 24.51 ATOM 207 C B LEU 700 30.274 0.397 -5.530 1.00 25.51 ATOM 208 CDL LEU 700 30.274 0.397 -5.537 1.00 19.67 ATOM 209 CDZ LEU 700 30.274 0.397 -5.537 1.00 19.67 ATOM 208 CDL LEU 700 30.274 0.397 -5.957 1.00 21.56 ATOM 209 CDZ LEU 700 30.274 0.397 -5.957 1.00 21.56 ATOM 208 CDL LEU 700 30.494 -7.957 1.00 21.56 ATOM 208 CDL LEU 700 30.494 -7.957 1.00 17.10 ATOM 210 C LEU 700 30.494 0.397 -5.957 1.00 17.10 ATOM 210 C LEU 700 30.495 -7.957 1.00 17.10 ATOM 210 C LEU 700 30.496 -7.977 1.00 17.57 ATOM 210 C LEU 700 30.496 -7.977 1.00 17.57 ATOM 210 C LEU 701 30.046 -2.991 -7.977 1.00 17.57 ATOM 210 C LEU 701 30.046 -2.991 -7.977 1.00 17.73 ATOM 210 C LEU 701 30.046 -2.991 -7.977 1.00 17.57 ATOM 210 C LEU 701 30.046 -2.991 -7.977 1.00 17.57 ATOM 210 C LEU 701 30.046 -2.991 -7.977 1.00 17.57 ATOM 210 C C B LEU 701						607	22 104	-3.669	-8.099	1.00 28.48
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ATOM 223 OG SER 702 37.204 0.120 -2.175 1.00 23.71 ATOM 224 C SER 702 34.874 2.791 -1.105 1.00 20.81 ATOM 225 O SER 702 35.187 3.423 -0.103 1.00 20.82 ATOM 226 N SER 703 34.023 3.250 -2.028 1.00 20.55 ATOM 227 CA SER 703 33.443 4.585 -1.934 1.00 18.91 ATOM 229 OG SER 703 32.755 4.966 -3.224 1.00 18.44 ATOM 229 OG SER 703 33.748 5.182 -4.194 1.00 20.63 ATOM 230 C SER 703 32.470 4.678 -0.793 1.00 18.35 ATOM 231 O SER 703 32.520 5.625 -0.025 1.00 18.89 ATOM 232 N LEU 704 31.596 3.684 -0.662 1.00 17.26 ATOM 233 CA LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 234 CB LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 235 CG LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 236 CD1 LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 236 CD1 LEU 704 27.882 1.259 -0.748 1.00 12.28 ATOM 237 CD2 LEU 704 27.582 3.681 -0.235 1.00 14.48 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 60 ATOM 242 CB ASN 705 33.3850 0.384 2.941 1.00 15.46 ATOM 242 CB ASN 705 33.850 0.384 2.941 1.00 16.284 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.68	40								-2.190	1.00 21.52
ATOM 224 C SER 702 34.874 2.791 -1.105 1.00 20.81 ATOM 225 O SER 702 35.187 3.423 -0.103 1.00 20.82 ATOM 226 N SER 703 34.023 3.250 -2.028 1.00 20.55 ATOM 227 CA SER 703 33.443 4.585 -1.934 1.00 18.91 ATOM 228 CB SER 703 32.755 4.966 -3.224 1.00 18.44 ATOM 229 OG SER 703 33.748 5.182 -4.194 1.00 20.63 ATOM 230 C SER 703 32.470 4.678 -0.793 1.00 18.35 ATOM 231 O SER 703 32.520 5.625 -0.025 1.00 18.89 ATOM 232 N LEU 704 31.596 3.684 -0.662 1.00 17.26 ATOM 233 CA LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 235 CG LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 235 CG LEU 704 28.558 2.583 -0.660 1.00 15.19 ATOM 236 CD1 LEU 704 27.882 1.259 -0.748 1.00 12.28 ATOM 237 CD2 LEU 704 27.882 1.259 -0.748 1.00 12.28 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 31.366 3.678 1.761 1.00 16.81 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.3850 0.384 2.941 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24	40									1.00 23.71
ATOM 225 O SER 702 35.187 3.423 -0.103 1.00 20.82 ATOM 226 N SER 703 34.023 3.250 -2.028 1.00 20.55 ATOM 227 CA SER 703 33.443 4.585 -1.934 1.00 18.91 ATOM 229 OG SER 703 32.755 4.966 -3.224 1.00 18.44 ATOM 229 OG SER 703 32.755 4.966 -3.224 1.00 18.44 ATOM 230 C SER 703 32.470 4.678 -0.793 1.00 18.35 ATOM 231 O SER 703 32.520 5.625 -0.025 1.00 18.89 ATOM 232 N LEU 704 31.596 3.684 -0.662 1.00 17.26 ATOM 233 CA LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 234 CB LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 235 CG LEU 704 28.558 2.583 -0.660 1.00 14.03 ATOM 236 CD1 LEU 704 27.582 1.259 -0.748 1.00 12.28 ATOM 237 CD2 LEU 704 27.582 3.681 -0.235 1.00 14.48 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.666 ATOM 242 CB ASN 705 33.850 0.384 2.941 1.00 16.24 ATOM 243 CG ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.28									-1.105	
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ATOM 230 C SER 703 32.470 4.678 -0.793 1.00 18.35 ATOM 231 O SER 703 32.520 5.625 -0.025 1.00 18.89 50 ATOM 232 N LEU 704 31.596 3.684 -0.662 1.00 17.26 ATOM 233 CA LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 234 CB LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 235 CG LEU 704 28.558 2.583 -0.660 1.00 14.03 ATOM 236 CD1 LEU 704 27.882 1.259 -0.748 1.00 12.28 55 ATOM 237 CD2 LEU 704 27.582 3.681 -0.235 1.00 14.48 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 60 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.28								5.182		
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50 ATOM 232 N LEU 704 31.596 3.684 -0.662 1.00 17.26 ATOM 233 CA LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 234 CB LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 235 CG LEU 704 28.558 2.583 -0.660 1.00 14.03 ATOM 236 CD1 LEU 704 27.882 1.259 -0.748 1.00 12.28 ATOM 237 CD2 LEU 704 27.582 3.681 -0.235 1.00 14.48 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.28								5.625	-0.025	
ATOM 233 CA LEU 704 30.639 3.687 0.432 1.00 16.37 ATOM 234 CB LEU 704 29.691 2.497 0.342 1.00 15.19 ATOM 235 CG LEU 704 28.558 2.583 -0.660 1.00 14.03 ATOM 236 CD1 LEU 704 27.882 1.259 -0.748 1.00 12.28 ATOM 237 CD2 LEU 704 27.582 3.681 -0.235 1.00 14.48 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.282	50					704		3.684		1.00 17.26
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ATOM 236 CD1 LEU 704 27.882 1.259 -0.748 1.00 12.28 ATOM 237 CD2 LEU 704 27.582 3.681 -0.235 1.00 14.48 ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24						704	28.558	2.583	-0.660	
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ATOM 238 C LEU 704 31.366 3.678 1.761 1.00 16.10 ATOM 239 O LEU 704 30.925 4.340 2.696 1.00 16.81 ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24	55							3.681		
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ATOM 240 N ASN 705 32.495 2.961 1.829 1.00 16.70 ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24										
ATOM 241 CA ASN 705 33.307 2.863 3.049 1.00 16.66 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24								2.961		
60 ATOM 242 CB ASN 705 34.398 1.794 2.924 1.00 15.46 ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24								2.863		
ATOM 243 CG ASN 705 33.850 0.384 2.941 1.00 16.24	60						34.398			
	-					705		0.384		
						705	34.448	-0.512	2.385	1.00 16.82

	ATOM	245	ND2	ASN	705	32.726	0.180	3.592	1.00 16.07
	ATOM	246	C	ASN	705	33.955	4.201	3.410	1.00 17.17
	ATOM	247	ŏ	ASN	705	33.970	4.587	4.570	1.00 17.46
	ATOM	248	N	GLU	706	34.512	4.882	2.415	1.00 17.04
5	ATOM	249	CA	GLU	706	35.151	6.193	2.598	1.00 17.55
J	ATOM	250	CB	GLU	706	35.739	6.668	1.258	1.00 18.93
		251	CG	GLU	706	36.394	8.029	1.282	1.00 21.19
	ATOM		CD	GLU	706	37.488	8.146	2.347	1.00 23.68
	ATOM	252			706	37.586	9.225	2.978	1.00 25.14
40	ATOM	253	OE1		706	38.246	7.175	2.569	1.00 24.37
10	ATOM	254	OE2			34.089	7.180	3.069	1.00 16.10
	ATOM	255	C	GLU	706		8.023	3.950	1.00 16.62
	MOTA	256	0	GLU	706	34.313		2.445	1.00 15.13
	ATOM	257	N	LEU	707	32.927	7.076		1.00 13.13
	ATOM	258	CA	LEU	707	31.803	7.916	2.792	1.00 14.21
15	ATOM	259	CB	LEU	707	• • •	7.579	1.925	
	ATOM	260	CG	LEU	707	29.318	8.262	2.328	1.00 12.03
	ATOM	261		LEU	707	29.537	9.745	2.280	1.00 13.09
	ATOM	262	CD2	LEU	707	28.252	7.889	1.374	1.00 12.74
	ATOM	263	С	LEU	707	31.461	7.634	4.228	1.00 14.91
20	ATOM	264	0	LEU	707	31.121	8.557	4.980	1.00 15.85
	ATOM	265	N	GLY	708	31.532	6.358	4.602	1.00 13.93
	ATOM	266	CA	GLY	708	31.230	5.976	5.965	1.00 12.96
	ATOM	267	С	GLY	708	32.213	6.620	6.917	1.00 13.13
	ATOM	268	0	GLY	708	31.849	7.061	7.987	1.00 13.55
25	ATOM	269	N	GLU	709	33.468	6.687	6.514	1.00 14.14
23	ATOM	270	CA	GLU	709	34.525	7.279	7.322	1.00 15.83
	ATOM	271	СВ	GLU	709	35.874	7.046	6.658	1.00 16.73
	ATOM	272	CG	GLU	709	37.051	7.547	7.446	1.00 18.68
	ATOM	273	CD	GLU	709	37.573	6.514	8.401	1.00 21.63
30	ATOM	274		GLU	709	36.766	5.660	8.826	1.00 23.39
30		275	OE2		709	38.784	6.544	8.723	1.00 23.17
	ATOM	276	C	GLU	709	34.334	8.775	7.486	1.00 16.65
	ATOM				709	34.628	9.317	8.563	1.00 17.59
	ATOM	277	0	GLU	710	33.845	9.427	6.428	1.00 16.70
25	ATOM	278	N	ARG	710	33.616	10.869	6.418	1.00 17.32
35	ATOM	279	CA	ARG		33.459	11.346	4.990	1.00 16.07
	ATOM	280	CB	ARG	710		11.098	4.137	1.00 16.18
	ATOM	281	CG	ARG	710	34.659	11.498	2.706	1.00 16.39
	ATOM	282	CD	ARG	710	34.329	11.436	1.850	1.00 15.28
40	ATOM	283	NE	ARG	710	35.512			1.00 15.20
40	ATOM	284	CZ	ARG	710	35.587	12.246	0.733	1.00 13.30
	ATOM	285		ARG	710	34.550	12.975	0.357	-
	ATOM	286		ARG	710	36.691	12.242	0.001	1.00 14.89
	ATOM	287	С	ARG	710	32.376	11.230	7.218	1.00 17.85
	ATOM	288	0	ARG	710	32.379	12.156	8.034	1.00 17.75
45	ATOM	289	N	GLN	711	31.291	10.516	6.955	1.00 18.71
	ATOM	290	CA	GLN	711	30.067	10.745	7.697	1.00 19.38
	ATOM	291	CB	GLN	711	28.908	9.938	7.127	1.00 19.79
	ATOM	292	CG	GLN	711	28.377	10.566	5.878	1.00 22.36
	ATOM	293	CD	GLN	711	27.058	10.010	5.446	1.00 23.37
50	ATOM	294	OE1	GLN	711	26.758	9.932	4.244	1.00 25.35
	ATOM	295	NE2	GLN	711	26.228	9.677	6.410	1.00 24.52
	ATOM	296	С	GLN	711	30.209	10.494	9.188	1.00 19.48
	ATOM	297	0	GLN	711	29.564	11.183	9.985	1.00 19.57
	ATOM	298	N	LEU	712	31.043	9.529	9.571	1.00 18.76
55	ATOM	299	CA	LEU	712	31.259	9.227	10.984	1.00 19.20
	ATOM	300	СВ	LEU	712	32.163	8.008	11.157	1.00 20.55
	ATOM	301	CG	LEU	712	32.522	7.607	12.590	1.00 21.95
	ATOM	302		LEU	712	31.288	7.641	13.484	1.00 23.43
	ATOM	303		LEU	712	33.132	6.223	12.586	1.00 22.57
60	ATOM	304	C	LEU	712	31.876	10.428	11.704	1.00 19.23
00		305	0	LEU	712	31.507	10.743	12.834	1.00 17.65
	ATOM				713	32.809	11.099	11.039	1.00 17.03
	ATOM	306	N	VAL	113	32.009	11.000	11.033	1.00 19.70

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		202	CD I	VAL	713	33.427	12.270	11.619	1.00 19.68
	ATOM	307 308		VAL	713	34.453	12.859	10.658	1.00 20.01
	ATOM ATOM	309		VAL	713	34.722	14.292	11.001	1.00 21.16
	MOTA	310		VAL	713	35.750	12.069	10.750	1.00 20.06
5	ATOM	311	C '	VAL	713	32.328	13.277	11.990	1.00 19.53 1.00 20.00
_	ATOM	312		VAL	713	32.325	13.802	13.086 11.128	1.00 20.00 1.00 19.20
	MOTA	313		HIS	714	31.330	13.434 14.356	11.128	1.00 19.20
	ATOM	314		HIS	714	30.215 29.498	14.558	10.038	1.00 20.77
40	ATOM	315		HIS HIS	714 714	30.331	15.410	9.058	1.00 21.60
10	ATOM	316	CG CD2		714	31.369	15.016	8.283	1.00 22.31
	ATOM ATOM	317 318	ND1		714	30.131	16.744	8.784	1.00 22.32
	ATOM	319	CEI		714	31.005	17.139	7.876	1.00 23.41
	ATOM	320	NE2		714	31.768	16.113	7.557	1.00 23.22
15	ATOM	321	С	HIS	714	29.183	13.885	12.383	1.00 18.83
	ATOM	322		HIS	714	28.497	14.701	13.005	1.00 18.73 1.00 18.39
	ATOM	323		VAL	715	29.006	12.572 11.972	12.485 13.434	1.00 16.86
	MOTA	324		VAL	715	28.063 27.869	10.435	13.134	1.00 16.78
00	ATOM	325	CB CG1	VAL	715 715	27.037	9.756	14.197	1.00 17.10
20	ATOM	326 327	CG1		715	27.183	10.259	11.817	1.00 17.34
	ATOM ATOM	328		VAL	715	28.667	12.166	14.817	1.00 15.60
	MOTA	329		VAL	715	27.958	12.422	15.788	1.00 15.49
	ATOM	330		VAL	716	29.986	12.077	14.913	1.00 15.13
25	ATOM	331	CA	VAL	716	30.622	12.250	16.205	1.00 15.01
	ATOM	332	CB	VAL	716	32.136	11.885	16.158 17.481	1.00 14.93 1.00 13.26
	ATOM	333		VAL	716	32.825 32.310	12.233 10.373	15.870	1.00 13.20
	ATOM	334	CG2		716 716	30.419	13.681	16.708	1.00 15.83
20	ATOM	335 336	C O	VAL VAL	716	30.129	13.883	17.887	1.00 16.61
30	ATOM ATOM	337	N	LYS	717	30.544	14.665	15.816	1.00 16.59
	ATOM	338	CA	LYS	717	30.390	16.082	16.183	1.00 17.20
	ATOM	339	СВ	LYS	717	30.884	16.974	15.041	1.00 18.94
	ATOM	340	CG	LYS	717	32.361	16.747	14.698	1.00 22.56
35	ATOM	341	CD	LYS	717	33.245	16.752	15.978 16.007	1.00 25.34 1.00 27.06
	MOTA	342	CE	LYS	717	34.294 34.709	15.609 15.195	17.410	1.00 27.00
	ATOM	343	NZ	LYS LYS	717 717	28.951	16.387	16.534	1.00 16.77
	ATOM ATOM	344 345	C O	LYS	717	28.658	16.931	17.593	1.00 18.49
40	ATOM	346	N	TRP	718	28.049	15.976	15.659	1.00 15.68
40	MOTA	347	CA	TRP	718	26.618	16.143	15.868	1.00 14.61
	ATOM	348	CB	TRP	718	25.889	15.442	14.689	1.00 11.97
	ATOM	349	CG	TRP	718	24.433	15.266	14.841	1.00 9.66
	ATOM	350	CD2	TRP	718	23.757	14.069	15.254	1.00 10.28 1.00 9.98
45	ATOM	351	CE2	TRP	718	22.373	14.371 12.778	15.293 15.612	1.00 10.09
	MOTA	352	CE3	TRP	718	24.176 23.472	16.199	14.645	1.00 9.89
	ATOM	353 354	CD1 NE1		718 718	22.228	15.688	14.918	1.00 8.38
	ATOM ATOM	355	CZ2		718	21.394	13.419	15.663	1.00 9.00
50	ATOM	356	CZ3		718	23.201	11.835	15.980	1.00 8.20
-	ATOM	357	CH2	TRP	718	21.835	12.171	16.004	1.00 7.32
	ATOM	358	С	TRP	718	26.200	15.562	17.261	1.00 15.34
	ATOM	359	0	TRP	718	25.659	16.269	18.124	1.00 14.55
	MOTA	360	N	ALA	719	26.468	14.272	17.464	1.00 16.10 1.00 15.03
55	ATOM	361	CA	ALA	719 719	26.143 26.796	13.559 12.184	18.683 18.657	1.00 13.03
	ATOM	362 363	CB C	ALA ALA	719 719	26.796	14.346	19.881	1.00 15.62
	ATOM ATOM	364	Ö	ALA	719	25.857	14.646	20.785	1.00 15.85
	ATOM	365	N	LYS	720	27.870	14.781	19.828	1.00 17.45
60	ATOM	366	CA	LYS	720	28.463	15.516	20.924	1.00 18.63
-	ATOM	. 367	СВ	LYS	720	29.970	15.625	20.715	1.00 19.81
	MOTA	368	CG	LYS	720	30.644	14.292	21.012	1.00 21.18

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	ATOM ATOM	369 CD 370 CE 371 NZ 372 C	LYS LYS LYS LYS	720 720 720 720	32.136 32.762 32.729 27.822	12.975 12.661 16.860	21.244 22.708 21.204	1.00 23.81 1.00 25.84 1.00 26.70 1.00 18.98 1.00 19.86
5	ATOM ATOM ATOM ATOM ATOM	373 O 374 N 375 CA 376 CB	LYS ALA ALA ALA	720 721 721 721	27.921 27.070 26.406 26.584 24.941	17.369 18.651	20.238 20.382 19.146 20.675	1.00 18.21 1.00 18.10 1.00 17.43 1.00 18.80
10	ATOM ATOM ATOM ATOM ATOM	377 C 378 O 379 N 380 CA 381 CB	ALA ALA LEU LEU LEU	721 721 722 722 722	24.192 24.518 23.119 22.955	19.485 17.247 16.912 15.395	20.660 20.904 21.207 21.119 20.271	1.00 19.16 1.00 19.00 1.00 19.60 1.00 19.45 1.00 19.62
15	ATOM ATOM ATOM ATOM	382 CG 383 CD	LEU LEU	722 722 722 722	21.855 21.540 22.298 22.754 23.521	14.771 15.657 13.382 17.362 17.125	19.099 19.815 22.616 23.549	1.00 17.02 1.00 17.38 1.00 20.27 1.00 21.72
20	ATOM ATOM ATOM ATOM ATOM	386 O 387 N 388 CD 389 CA 390 CB	PRO	722 723 723 723 723 723	21.574 20.500 21.211 19.767 19.706	17.992 18.317 18.428 18.917 19.349	22.811 21.861 24.167 23.997 22.624	1.00 20.69 1.00 20.29 1.00 21.24 1.00 20.40 1.00 20.05
25	ATOM ATOM ATOM ATOM ATOM	391 CG 392 C 393 O 394 N 395 CA 396 C	PRO PRO GLY	723 723 724 724 724	21.266 20.821 21.800 21.874 22.838	17.287 16.165 17.588 16.598 15.478	25.195 24.935 26.369 27.416 27.132	1.00 21.66 1.00 21.14 1.00 22.02 1.00 22.29 1.00 23.13 1.00 23.78
30	ATOM ATOM ATOM ATOM ATOM	397 O 398 N 399 CA 400 CE 401 CO	GLY PHE PHE PHE	724 725 725 725 725 725	23.076 23.434 24.360 24.915 25.648	14.554 13.353	28.004 25.946 25.610 24.214 23.703	1.00 24.14 1.00 24.24 1.00 23.59 1.00 23.80 1.00 22.83
35	MOTA MOTA MOTA MOTA MOTA	402 CE 403 CE 404 CE 405 CE	D1 PHE D2 PHE E1 PHE E2 PHE	725 725 725 725 725 725	24.944 27.046 25.623 27.731 27.025	13.328 11.130 12.226	23.260 23.675 22.804 23.221 22.784	1.00 22.40 1.00 22.77 1.00 21.05 1.00 22.31
40	ATOM ATOM ATOM ATOM	406 C3 407 C 408 O 409 N 410 C 411 C	PHE PHE ARG A ARG	725 725 725 726 726 726	25.505 25.873 26.083 27.207 27.831	14.170 13.028 15.270 15.229 16.620	26.582 26.863 27.070 28.033 28.204	1.00 24.85 1.00 23.79 1.00 25.97 1.00 27.63 1.00 29.27
45	ATOM ATOM ATOM ATOM ATOM ATOM	412 C 413 C 414 N 415 C	G ARG D ARG E ARG	726 726 726 726 726 726	28.622 29.759 30.65 31.873 32.63	9 16.141 7 16.595 2 16.090 5 16.558	25.670 25.464 24.486	1.00 31.68 1.00 34.22 1.00 37.18 1.00 38.28 1.00 39.44 1.00 38.78
50	ATOM ATOM ATOM ATOM	417 N 418 C 419 O 420 N	H2 ARG ARG ARG I ASN	726 726 726 727 727	32.32 26.90 27.79 25.63 25.24	2 14.615 7 14.449 2 14.316	29.423 30.253 29.683	1.00 27.24 1.00 27.08 1.00 27.15 1.00 26.29
55	MOTA MOTA MOTA MOTA MOTA	422 C 423 C 424 C 425 N	B ASN G ASN DD1 ASN ND2 ASN	727 727 727 727	23.71 23.11 21.94 23.90 25.75	7 13.687 8 15.085 7 15.277 9 16.055	31.115 31.195 30.893 31.628	1.00 24.95 1.00 24.88 1.00 24.54 1.00 26.17
60	ATOM ATOM ATOM ATOM ATOM	427 0 428 1 429 0	ASN ASN LEU CA LEU CB LEU	727 728 728	25.75 25.99 25.89 26.36	12 11.689 15 11.67 15 10.28	31.994 3 29.749 9 29.629	1.00 27.30 1.00 25.64 1.00 25.16

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1.00 20.68 9.711 27.448 24.859 728 CG LEU 1.00 19.66 431 26.076 9.133 MOTA 25.084 CD1 LEU 728 1.00 19.79 432 28.203 ATOM 23.856 8.883 728 CD2 LEU 1.00 25.79 433 10.208 29.974 ATOM 27.833 LEU 728 1.00 25.15 С 434 29.739 11.157 ATOM 28.571 728 1.00 27.05 LEU 0 435 30.516 ATOM 9.064 28.247 729 1.00 28.68 HIS 436 N 30.871 ATOM 8.808 29.642 729 HIS 1.00 30.72 CA 31.570 437 MOTA 7.455 29.737 729 438 HIS 1.00 33.13 CB 31.943 ATOM 7.042 31.132 729 1.00 33.50 HIS 31.218 439 CG MOTA 32.276 6.978 .729 1.00 34.21 CD2 HIS 440 33.209 10 MOTA 6.603 31.460 729 ND1 HIS 1.00 34.64 33.247 441 ATOM 32.744 6.293 729 1.00 34.52 CE1 HIS 442 32.049 ATOM 6.510 33.263 729 1.00 29.25 1.00 29.28 NE2 HIS 443 29.577 MOTA 8.772 30.450 729 HIS 444 С 28.594 MOTA 8.182 30.003 729 HIS 1.00 30.21 0 445 9.295 15 MOTA 29.625 31.681 730 VAL 1.00 31.02 446 N 28.465 MOTA 9.365 32.592 730 CA VAL 1.00 32.08 447 28.898 MOTA 34.036 9.793 CB VAL 730 1.00 32.91 448 MOTA 9.446 27.811 35.077 CG1 VAL 730 1.00 31.86 29.176 449 MOTA 11.284 34.074 730 1.00 30.89 450 CG2 VAL 27.600 20 MOTA 8.108 32.662 730 VAL 1.00 30.40 451 C 26.371 MOTA 8.192 32.704 730 VAL 1.00 30.81 452 0 28.244 MOTA 32.770 6.956 27.509 1.00 31.55 28.410 1.00 36.23 27.771 1.00 40.32 731 453 N ASP MOTA 5.709 32.819 CA ASP 731 454 MOTA 4.536 33.244 731 CB ASP 455 25 MOTA 3.152 32.966 456 CG ASP 731 27.974 1.00 42.21 MOTA 2.619 31.837 731 2.599 27.075 1.00 42.23 OD1 ASP MOTA 457 33.867 731 458 OD2 ASP 1.00 28.94 5.425 26.889 ATOM 31.474 731 ASP 1.00 29.47 4.912 25.789 MOTA 459 С 31.408 731 ASP 1.00 26.17 460 0 5.760 27.587 30 MOTA 30.403 732 ASP 461 N 27.057 1.00 24.53 5.510 MOTA 29.079 732 ASP 1.00 24.04 CA 462 5.711 28.119 ATOM 28.024 ASP 732 1.00 23.64 1.00 22.31 463 CB 4.654 29.186 MOTA 28.073 CG ASP 732 464 MOTA 28.984 3.592 28.728 732 1.00 23.89 OD1 ASP 35 465 ATOM 30.231 4.904 27.444 OD2 ASP 732 1.00 23.96 466 25.875 MOTA 6.387 28.770 732 ASP 1.00 22.56 С 467 5.982 24.981 MOTA 28.030 ASP 732 1.00 23.56 468 0 25.920 MOTA 7.612 29.288 733 GLN N 24.855 1.00 22.94 469 ATOM 8.591 29.121 CA GLN 733 1.00 22.73 470 40 MOTA 25.166 9.847 29.942 733 CB GLN 1.00 23.24 471 MOTA 26.225 29.359 10.776 733 1.00 23.27 CG GLN 472 MOTA 26.480 12.013 30.208 733 GLN CD 1.00 24.33 1.00 23.47 473 27.477 MOTA 12.696 30.018 733 OE1 GLN 474 **ATOM** 25.577 31.130 29.636 28.979 12.316 733 NE2 GLN 1.00 23.20 475 45 MOTA 7.997 23.557 733 476 C GLN 1.00 23.08 ATOM 22.522 8.075 733 GLN 0 477 1.00 23.37 MOTA 7.459 23.625 30.853 30.00 734 478 N MET MOTA 22.508 1.00 24.31 6.832 734 CA MET 1.00 27.26 479 MOTA 22.906 6.596 33.003 480 CB MET 734 1.00 31.61 50 MOTA 22.047 5.604 33.749 734 481 CG MET 1.00 39.54 MOTA 22.821 35.293 5.121 SD MET 734 MOTA 482 1.00 37.92 23.387 3.401 34.884 483 CE MET 734 22.077 MOTA 1.00 23.31 5.510 30.902 734 MET 484 C 1.00 23.35 MOTA 20.884 5.247 30.732 485 O MET 734 55 1.00 21.64 ATOM 23.052 4.671 30.571 735 486 N ALA 1.00 20.44 ATOM 22.788 3.390 29.939 735 ALA 487 CA 1.00 20.71 MOTA 24.110 2.683 735 29.650 ALA 488 CB ATOM 1.00 19.23 22.013 28.644 3.570 735 ALA 1.00 19.59 С 489 MOTA 21.015 2.905 /36 27.799 736 26 51 735 28.398 ALA 0 490 1.00 18.69 60 MOTA 22.501 4.460 VAL 491 N 1.00 17.76 MOTA 21.877 4.734 492 CA VAL MOTA

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1.00 18.30 22.760 5.771 25.742 1.00 17.05 CB VAL 736 493 22,011 6.998 MOTA 25.373 736 1.00 17.25 CG1 VAL 494 5.118 23.420 MOTA 24.544 CG2 VAL 736 1.00 18.16 495 20.389 MOTA 5.133 26.673 736 1.00 17.42 VAL С 496 19.512 4.614 MOTA 25.962 1.00 17.60 736 VAL 20.096 497 0 5.985 ATOM 27.658 737 ILE 1.00 16.31 498 N 18.724 6.429 MOTA 27.914 737 1.00 14.71 ILE 18.683 CA MOTA 499 7.497 29.046 737 ILE 1.00 14.05 17.272 CB 500 ATOM 29.476 7.772 737 1.00 15.11 CG2 ILE 19.325 501 8.819 28.602 MOTA 1.00 12.32 737 CG1 ILE 502 19.618 10 MOTA 29.769 9.804 1.00 16.74 CD1 ILE 737 503 5.216 17.904 MOTA 28.352 737 1.00 16.09 ILE 504 С 16.812 MOTA 4.982 27.853 1.00 18.15 737 ILE 505 0 18.468 29.281 4.451 ATOM . 738 GLN 1.00 17.97 N 506 17.845 MOTA 3.260 29.850 738 1.00 20.53 GLN 507 CA 18.713 MOTA 2.715 15 30.960 1.00 23.88 GLN 738 CB 508 18.568 32.278 3.394 MOTA 738 1.00 26.69 CG GLN 19.439 509 2.726 MOTA 33.306 GLN 738 1.00 29.30 20.593 510 CD ATOM 2.390 33.027 738 1.00 28.53 OE1 GLN 511 18.887 MOTA 2.475 34.483 738 1.00 16.76 NE2 GLN 512 17.548 20 ATOM 2.111 28.904 738 1.00 16.47 С GLN 513 16.788 1.226 29.249 MOTA 738 1.00 16.39 GLN 0 514 2.029 18.260 MOTA 27.792 739 1.00 16.41 TYR 17.995 515 N MOTA 26.819 0.983 739 CA TYR 1.00 15.99 MOTA 516 19.285 0.448 26.174 1.00 15.68 739 TYR 20.313 517 CB 25 ATOM -0.115 27.130 739 TYR 1.00 15.41 CG 518 19.950 MOTA -0.852 28.251 739 CD1 TYR 1.00 16.17 519 20.915 MOTA -1.31729.151 CE1 TYR 739 1.00 16.49 520 21.656 MOTA 0.131 26.925 739 CD2 TYR 1.00 17.09 22.624 521 **ATOM** -0.321 27.817 739 CE2 TYR 1.00 16.83 522 -1.040 22.253 30 MOTA 28.921 23.256 739 1.00 18.24 TYR CZ 523 MOTA -1.435 29.787 739 TYR 524 OH 17.100 1.00 16.45 1.527 MOTA 25.721 1.00 17.63 TYR 739 С 16.295 525 MOTA 0.793 25.138 739 TYR 1.00 15.61 526 0 17.195 ATOM 2.822 25.453 SER 740 1.00 15.61 527 N 16.403 35 MOTA 3.404 24.384 740 1.00 15.49 CA SER 528 4.403 17.252 MOTA 23.619 740 SER 1.00 18.39 CB 529 17.682 ATOM 5.421 24.512 740 SER 1.00 14.87 OG 530 15.060 MOTA 4.054 24.697 740 1.00 15.75 531 С SER MOTA 14.376 4.451 23.778 740 1.00 14.33 SER 0 MOTA 532 14.659 40 25.948 4.188 741 1.00 14.83 TRP N 533 MOTA 4.835 13.382 26.202 741 TRP 1.00 15.35 534 CA 27.706 13.113 MOTA 4.997 741 TRP CB 1.00 17.05 535 MOTA 13.000 3.720 28.465 741 CG TRP 11.800 1.00 17.91 536 2.987 MOTA 28.765 CD2 TRP 741 1.00 18.98 45 ATOM 537 1.834 12.190 29.467 741 CE2 TRP 1.00 18.66 538 ATOM 10.434 3.193 28.505 CE3 TRP 741 1.00 17.12 MOTA 539 14.020 28.995 3.016 741 1.00 19.60 540 CD1 TRP MOTA 13.551 1.878 29.592 NE1 TRP 741 1.00 18.79 541 11.266 0.876 MOTA 29.915 741 CZ2 TRP 1.00 17.91 50 542 9.509 MOTA 28.949 2.240 741 CZ3 TRP 543 1.00 18.35 MOTA 9.934 1.098 29.644 741 CH2 TRP 1.00 15.12 544 MOTA 12.166 25.471 4.246 741 545 C TRP 1.00 15.56 4.995 MOTA 11.376 24.902 741 TRP 0 12.034 546 1.00 14.55 MOTA 2.920 25.391 742 MET 1.00 12.90 Ν 55 547 **ATOM** 2.339 10.870 24.723 742 CA MET 1.00 13.16 548 MOTA 10.866 0.815 24.785 742 MET CB 1.00 12.17 549 MOTA 9.597 24.219 0.185 742 MET CG 1.00 15.00 550 MOTA 0.336 8.263 25.353 742 MET SD 551 1.00 13.52 MOTA 8.639 26.462 -0.994 742 MET 1.00 12.47 552 CE 60 MOTA 2.763 10.699 23.290 742 MET 9.610 1.00 13.31 553 С MOTA 3.153 22.886 742 MET 554 0 MOTA



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					7.40	22 407	2 656	11.748	1.00 12.12
	MOTA	555	-	GLY	743	22.497	2.656	11.748	1.00 11.81
	ATOM	556		GLY	743	21.102	3.057		1.00 12.08
	ATOM	557	_	GLY	743	20.947	4.558	11.452 10.768	1.00 11.88
	ATOM	558	-	GLY	743	20.022	5.009	·	1.00 12.77
5	ATOM	559		LEU	744	21.835	5.336	12.070	1.00 12.77
	ATOM	560		LEU	744	21.817	6.797	11.972 12.888	1.00 12.22
	ATOM	561	-	LEU	744	22.884	7.418		1.00 11.70
	ATOM	562		LEU	744	22.702	7.481	14.399	1.00 9.77
	ATOM	563	CD1		744	23.967	8.075	14.954	1.00 9.16
10	MOTA	564	CD2		744	21.516	8.341	14.799	1.00 12.20
	ATOM	565		LEU	744	22.087	7.258	10.563	1.00 12.20
	ATOM	566		LEU	744	21.424	8.173	10.080	1.00 14.22
	ATOM	567		MET	745	23.083	6.651	9.921	1.00 12.34
	MOTA	568		MET	745	23.466	6.991	8.541	1.00 10.75
15	ATOM	569		MET	745	24.839	6.427	8.191	
	ATOM	570	CG	MET	745	25.961	6.948	9.076	
	MOTA	571	-	MET	745	27.509	6.429	8.487	
	MOTA	572	CE	MET	745	28.579	6.939	9.717	1.00 9.84 1.00 12.29
	ATOM	573	-	MET	745	22.462	6.498	7.508	
20	MOTA	574		MET	745	22.234	7.155	6.495	1.00 10.62
	ATOM	575	N	VAL	746	21.855	5.342	7.793	1.00 12.05
	ATOM	576		VAL	746	20.874	4.733	6.934	1.00 11.50
	ATOM	577		VAL	746	20.524	3.315	7.426	1.00 11.19
	ATOM	578	CG1	VAL	746	19.245	2.852	6.811	1.00 10.17
25	ATOM	579	CG2	VAL	746	21.615	2.355	7.095	1.00 9.64
	ATOM	580	С	VAL	746	19.605	5.565	6.942	1.00 12.13
	ATOM	581	0	VAL	746	19.000	5.792	5.907	1.00 12.72
	ATOM	582	N	PHE	747	19.227	6.051	8.117	1.00 12.64
	ATOM	583	CA	PHE	747	18.014	6.857	8.304	1.00 12.63
30	ATOM	584	CB	PHE	747	17.763	7.031	9.800	1.00 11.19
	ATOM	585	CG	PHE	747	16.411	7.542	10.126	1.00 10.00
	ATOM	586	CD1	PHE	747	15.286	6.780	9.847	1.00 9.30
	ATOM	587	CD2		747	16.253	8.798	10.700	1.00 7.79
	ATOM	588	CE1	PHE	747	14.008	7.260	10.136	1.00 8.30
35	ATOM	589	CE2		747	14.996	9.293	10.993	1.00 6.75
	ATOM	590	ÇZ	PHE	747	13.867	8.524	10.707	1.00 8.21
	ATOM	591	С	PHE	747	18.137	8.241	7.621	1.00 13.26
	ATOM	592	0	PHE	747	17.178	8.751	7.042	1.00 13.81
	ATOM	593	N	ALA	748	19.298	8.873	7.740	1.00 12.46
40	ATOM	594	CA	ALA	748	19.513	10.172	7.119	1.00 12.97
	ATOM	595	CB	ALA	748	20.749	10.808	7.648	1.00 11.50
	ATOM	596	С	ALA	748	19.640	9.988	5.635	1.00 13.78
	MOTA	597	0	ALA	748	19.226	10.850	4.882	1.00 14.44
	ATOM	598	N	MET	749	20.209	8.864	5.204	1.00 14.54
45	ATOM	599	CA	MET	749	20.381	8.578	3.782	1.00 14.78 1.00 15.28
	ATOM	600	СВ	MET	749	21.241	7.331	3.607	
	MOTA	601	CG	MET	749	21.622	6.945	2.199	1.00 15.33
	MOTA	602	SD	MET	749	20.315	6.246	1.193	1.00 18.79
	ATOM	603	CE	MET	749	20.226	4.627	1.835	1.00 18.82
50	ATOM	604	С	MET	749	19.023	8.390	3.142	1.00 15.85 1.00 17.51
	ATOM	605	0	MET	749	18.808	8.780	1.990	
	ATOM	606	N	GLY	750	18.088	7.829	3.895	1.00 16.02
	ATOM	607	CA	GLY	750	16.748	7.618	3.384	1.00 16.34
	ATOM	608	С	GLY	750	16.057	8.956	3.225	1.00 17.79
55	ATOM	609	0	GLY	750	15.263	9.135	2.289	1.00 19.00
	ATOM	610	N	TRP	751	16.361	9.897	4.121	1.00 17.36
	MOTA	611	CA	TRP	751	15.778	11.241	4.091	1.00 17.99
	ATOM	612	СВ	TRP	751	16.108	12.026	5.366	1.00 16.08
	ATOM	613	CG	TRP	751	15.528	13.458	5.416	1.00 14.99
60	ATOM	614		TRP	751	14.151	13.821	5.617	1.00 13.68
	ATOM	615		TRP	751	14.099	15.230	5.697	1.00 12.98
	ATOM	616	CE3	TRP	751	12.967	13.090	5.743	1.00 14.14

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								5 364	1.00 13.27
	ATOM	617	CD1	TRP	751	16.225	14.636	5.364	1.00 13.27
	ATOM	618	NE1	TRP	751	15.375	15.705	5.538	1.00 12.27
	ATOM	619	CZ2	TRP	751	12.907	15.926	5.899	1.00 14.74
	ATOM	620	CZ3	TRP	751	11.775	13.780	5.942	
5	ATOM	621	CH2	TRP	751	11.756	15.188	6.020	
•	ATOM	622	С	TRP	751	16.266	11.995	2.857	
	ATOM	623	0	TRP	751	15.457	12.558	2.124	1.00 20.13
	ATOM	624	N	ARG	752	17.569	11.971	2.607	1.00 19.13
	ATOM	625	CA	ARG	752	18.150	12.616	1.431	1.00 19.06
10	ATOM	626	CB	ARG	752	19.644	12.380	1.389	1.00 18.53
. •	ATOM	627	CG	ARG	752	20.370	12.908	2.567	1.00 18.25
	ATOM	628	CD	ARG	752	21.870	12.901	2.317	1.00 17.24
	ATOM	629	NE	ARG	752	22.467	11.573	2.298	1.00 14.94
	ATOM	630	CZ	ARG	752	22.976	10.973	3.370	1.00 14.90
15	ATOM	631	NH1	ARG	752	22.928	11.561	4.554	1.00 14.75
	ATOM	632	NH2	ARG	752	23.684	9.864	3.240	1.00 13.87
	ATOM	633	С	ARG	752	17.572	12.077	0.138	1.00 20.27 1.00 20.66
	ATOM	634	0	ARG	752	17.392	12.815	-0.828	1.00 20.66 1.00 22.00
	ATOM	635	N	SER	753	17.391	10.761	0.083	1.00 22.00
20	ATOM	636	CA	SER	753	16.823	10.099	-1.093	1.00 22.25
	ATOM	637	CB	SER	753	16.716	8.590	-0.879 -0.687	1.00 20.23
	MOTA	638	OG	SER	753	17.988	8.027		1.00 23.31
	ATOM	639	С	SER	753	15.434	10.635	-1.289	1.00 23.88
	ATOM	640	0	SER	753	14.978	10.803	-2.409 -0.173	1.00 23.00
25	ATOM	641	N	PHE	754	14.762	10.870	-0.173	1.00 24.70
	ATOM	642	CA	PHE	754	13.405	11.375	1.245	1.00 26.43
	ATOM	643	CB	PHE	754	12.835	11.243 11.765	1.364	1.00 28.06
	MOTA	644	CG	PHE	754	11.447		0.654	1.00 28.69
	MOTA	645	CD1	PHE	754	10.407	11.168	2.118	1.00 27.96
30	ATOM	646	CD2	PHE	754	11.184 9.126	12.895 11.703	0.687	1.00 29.47
	MOTA	647	CE1		754		13.442	2.160	1.00 28.93
	ATOM	648	CE2	PHE	754	9.901 8.876	12.849	1.445	1.00 29.47
	ATOM	649	CZ	PHE	754	13.239	12.818	-0.630	1.00 27.47
	ATOM	650	C	PHE	754 754	12.543	13.100	-1.614	1.00 26.87
35	MOTA	651	0	PHE	754 755	13.823	13.732	0.125	1.00 29.01
	MOTA	652	N	THR	755 755	13.725	15.134	-0.190	1.00 30.83
	ATOM	653	CA	THR	755 755	14.345	15.972	0.918	1.00 29.71
	ATOM	654	CB	THR THR	755 755	15.669	15.524	1.183	1.00 28.99
40	ATOM	655 656	OG1 CG2		755	13.553	15.796	2.164	1.00 29.63
40	ATOM	657	CGZ	THR	755	14.317	15.460	-1.552	1.00 32.57
	ATOM	658	0	THR	755	13.841	16.358	-2.234	1.00 33.24
	ATOM	659	N	ASN	756	15.262	14.639	-1.991	1.00 34.71
	ATOM ATOM	660	CA	ASN	756	15.920	14.842	-3.273	1.00 36.48
45	ATOM	661	CB	ASN	756	17.417	14.562	-3.149	1.00 36.89
40	ATOM	662	CG	ASN	756	18.137	15.616	-2.344	1.00 37.02
	ATOM	663		ASN	756	17.563	16.237	-1.456	1.00 39.11
	ATOM	664		ASN	756	19.392	15.844	-2.668	1.00 37.24
	ATOM	665	C	ASN	756	15.360	14.065	-4.457	1.00 37.88
50	ATOM	666	ŏ	ASN	756	14.684	14.628	-5.313	1.00 39.57
••	ATOM	667	N	VAL	757	15.654	12.773	-4.518	1.00 38.99
	ATOM	668	CA	VAL	757	15.210	11.948	-5.633	1.00 39.74
	ATOM	669	СВ	VAL	757	16.274	10.869	-5.971	1.00 39.96
	ATOM	670	CG1		757	17.639	11.540	-6.170	1.00 40.00
55	ATOM	671		VAL	757	16.354	9.819	-4.871	1.00 39.28
- •	ATOM	672	С	VAL	757	13.835	11.308	-5.456	1.00 39.95
	ATOM	673	ŏ	VAL	757	13.501	10.335	-6.134	1.00 40.19
	ATOM	674	N	ASN	758	13.037	11.874	-4.559	1.00 40.46
	ATOM	675	CA	ASN	758	11.699	11.374	-4.265	1.00 41.28
60	ATOM	676	СВ	ASN	758	10.678	11.894	-5.288	1.00 43.82
	ATOM	677	CG	ASN	758	10.257	13.331	-5.005	1.00 44.84
	MOTA	678	OD1	ASN	758	11.097	14.199	-4.764	1.00 46.40

	MOTA	679	ND2	ASN	758	8.953	13.576	-4.987	1.00	
	ATOM	680		ASN	758	11.622	9.858	-4.100		40.91
	ATOM	681		ASN	758	10.592	9.229	-4.404		40.73
	ATOM	682		SER	759	12.733	9.298	-3.612	1.00	
5	ATOM	683		SER	759	12.891	7.877	-3.326		38.71
5		684		SER	759	11.763	7.415	-2.395		37.53
	MOTA			SER	759	11.496	8.369	-1.378	1.00	34.26 .
	ATOM	685			759	13.027	6.921	-4.532	1.00	39.24
	MOTA	686		SER		12.833	5.711	-4.382	1.00	
	ATOM	687	-	SER	759	13.409	7.438	-5.704		39.12
10	MOTA	688		ARG	760			-6.892		38.62
	ATOM	689		ARG	760	13.564	6.589		1.00	-
	ATOM	690	CB	ARG	760	13.451	7.422	-8.171		
	ATOM	691	CG	ARG	760	13.598	6.577	-9.444		44.41
	ATOM	692	CD	ARG	760	13.903		-10.715	1.00	
15	ATOM	693	NE	ARG	760	14.534	6.544	-11.729		48.86
. •	ATOM	694		ARG	760	13.875	5.797	-12.614	1.00	49.74
	ATOM	695	NH1		760	12.542	5.795	-12.649	1.00	50.04
		696	NH2		760	14.553	4.969	-13.398	1.00	49.46
	ATOM		C	ARG	760	14.897	5.840	-6.876	1.00	36.88
20	ATOM	697		ARG	760	15.024	4.741	-7.426	1.00	37.48
20	ATOM	698	0			15.902	6.466	-6.275		34.87
	ATOM	699	N	MET	761	17.238	5.890	-6.159		32.21
	ATOM	700	CA	MET	761		6.510	-7.194		33.77
	MOTA	701	CB	MET	761	18.171		-8.571		36.10
	ATOM	702	CG	MET	761	17.588	6.682			40.36
25	ATOM	703	SD	MET	761	18.859	7.115	-9.788		
	ATOM	704	CE	MET	761	18.737	8.904	-9.809		38.10
	ATOM	705	С	MET	761	17.738	6.242	-4.751		29.46
	ATOM	706	0	MET	761	17:144	7.080	-4.075		28.57
	ATOM	707	N	LEU	762	18.837	5.635	-4.319		26.78
30	ATOM	708	CA	LEU	762	19.382	5.905	-2.992	1.00	24.13
30	ATOM	709	CB	LEU	762	19.956	4.637	-2.393		24.05
		710	CG	LEU	762	18.957	3.502	-2.272	1.00	23.69
	MOTA			LEU	762	19.615	2.272	-1.632	1.00	23.99
	MOTA	711			762	17.788	4.011	-1.439		24.34
0.5	ATOM	712		LEU		20.458	6.968	-3.040		23.01
35	MOTA	713	C	LEU	762		6.726	-3.548		22.65
	MOTA	714	0	LEU	762	21.537		-2.475		22.03
	ATOM	715	N	TYR	763	20.162	8.132			20.69
	ATOM	716	CA	TYR	763	21.066	9.273	-2.450	1.00	23.12
	ATOM	717	CB	TYR	763	20.250	10.540	-2.266		
40	ATOM	718	ÇG	TYR	763	20.946	11.782	-2.730		25.58
	ATOM	719	CD1	TYR	763	20.841	12.187	-4.052		26.87
	ATOM	720	CE1	TYR	763	21.416	13.373	-4.492	1.00	
	ATOM	721	CD2	TYR	763	21.662	12.590	-1.841	1.00	26.77
	ATOM	722	CE2		763	22.247	13.789	-2.272	1.00	28.35
45	ATOM	723	CZ	TYR	763	22.107	14.172	-3.604	1.00	28.85
70		724	ОН	TYR	763	22.595	15.379	-4.047	1.00	30.59
	ATOM	725	C	TYR	763	22.068	9.173	-1.323	1.00	18.78
	ATOM				763	21.910	9.828	-0.304		17.73
	ATOM	726	0	TYR		23.128	8.401	-1.538		17.33
	ATOM	727	N	PHE	764		8.191			16.94
50	ATOM	728	CA	PHE	764	24.152				15.91
	ATOM	729	CB	PHE	764	25.086	7.078			
	ATOM	730	CG	PHE	764	24.505	5.724			16.79
	ATOM	731		PHE	764	24.211	4.961			16.06
	MOTA	732	CD2	PHE	764	24.267	5.205	0.450	1.00	
55	ATOM	733		PHE	764	23.691	3.692			18.06
	ATOM	734		PHE	764	23.748	3.941	0.606	1.00	18.27
	ATOM	735	CZ	PHE	764	23.458	3.176			17.80
				PHE	764	24.964	9.441			17.39
	MOTA	736	C		764	25.379	9.797			17.28
00	ATOM	737	0	PHE		25.224	10.084			17.00
60	ATOM	738	N	ALA	765					16.32
	ATOM	739	CA	ALA	765	26.013	11.292			
	ATOM	740	СВ	ALA	765	27.479	10.957	-1.460	1.00	16.17

								2 243	1.00 16.66
	ATOM	741	C Z	ALA	765	25.674	11.913	-2.841	1.00 16.30
	ATOM	742	0 7	ALA	765	25.051	11.267	-3.675	
	ATOM	743	N	PRO	766	26.016	13.196	-3.032	1.00 17.18
	ATOM	744	CD	PRO	766	26.544	14.169	-2.064	1.00 15.41
5	ATOM	745	CA	PRO	766	25.703	13.846	-4.311	1.00 17.49
J	ATOM	746		PRO	766	26.183	15.277	-4.077	1.00 17.30
		747	-	PRO	766	26.002	15.451	-2.608	1.00 17.07
	ATOM	748		PRO	766	26.429	13.161	-5.481	1.00 17.65
	ATOM	749		PRO	766	25.923	13.099	-6.598	1.00 17.73
40	MOTA		-	ASP	767	27.578	12.569	-5.166	1.00 18.27
10	ATOM	750		ASP	767	28.416	11.850	-6.115	1.00 18.49
	MOTA	751				29.877	12.312	-5.955	1.00 18.71
	ATOM	752	-	ASP	767		12.135	-4.525	1.00 19.47
	ATOM	753		ASP	767	30.413		-3.569	1.00 20.31
	ATOM	754	OD1		767	29.611	12.038		1.00 19.04
15	ATOM	755	OD2		767	31.650	12.102	-4.348	
	ATOM	756	С	ASP	767	28.330	10.317	-5.981	1.00 17.79
	ATOM	757	0	ASP	767	29.191	9.594	-6.476	1.00 18.69
	ATOM	758	N	LEU	768	27.334	9.820	-5.267	1.00 18.04
	ATOM	759	CA	LEU	768	27.164	8.379	-5.110	1.00 18.20
20	ATOM	760		LEU	768	27.955	7.809	-3.914	1.00 17.47
20		761		LEU	768	28.032	6.263	-3.786	1.00 16.12
	MOTA	762	CD1		768	28.641	5.671	-5.047	1.00 14.30
	MOTA		CD2		768	28.850	5.846	-2.563	1.00 15.17
	ATOM	763			768	25.690	8.129	-4.930	1.00 18.58
	ATOM	764		LEU		25.184	8.068	-3.812	1.00 17.79
25	ATOM	765		LEU	768		8.156	-6.048	1.00 19.79
	MOTA	766	N	VAL	769	24.979		~6.035	1.00 20.42
	ATOM	767	CA	VAL	769	23.553	7.895		1.00 20.42
	ATOM	768	CB	VAL	769	22.709	9.142	-6.447	
	ATOM	769	CG1	VAL	769	23.571	10.190	-7.096	1.00 20.70
30	MOTA	770	CG2	VAL	769	21.537	8.757	-7.277	1.00 19.19
•	ATOM	771	С	VAL	769	23.373	6.609	-6.852	1.00 20.73
	ATOM	772	0	VAL	769	23.873	6.467	-7.961	1.00 22.43
	ATOM	773	N	PHE	770	22.871	5.604	-6.157	1.00 19.70
	ATOM	774	CA	PHE	770	22.683	4.277	-6.681	1.00 19.29
35	ATOM	775	СВ	PHE	770	22.596	3.263	-5.503	1.00 18.44
55	ATOM	776	CG	PHE	770	23.930	2.757	-4.996	1.00 16.41
	MOTA	777	CD1		770	25.079	3.546	-5.053	1.00 14.52
		778		PHE	770	24.025	1.468	-4.459	1.00 14.96
	ATOM			PHE	770	26.291	3.070	-4.588	1.00 13.39
40	ATOM	779		PHE	770	25.243	0.979	-3,983	1.00 13.90
40	MOTA	780	CE2		770	26.383	1.786	-4.050	1.00 13.96
	ATOM	781	CZ	PHE		_	4.134	-7.473	1.00 19.41
	ATOM	782	С	PHE	770	21.425		-7.054	1.00 19.74
	MOTA	783	0	PHE	770	20.367	4.583		1.00 20.33
	ATOM	784	N	ASN	771	21.534	3.474	-8.611	
45	ATOM	785	CA	ASN	771	20.363	3.157	-9.410	
	ATOM	786	CB	ASN	771	20.524		-10.864	1.00 19.33
	ATOM	787	CG	ASN	771	21.883		-11.403	1.00 18.89
	ATOM	788	OD1	ASN	771	22.574		-10.942	1.00 19.51
	ATOM	789		ASN	771	22.289	4.069	-12.382	1.00 19.02
50	ATOM	790	С	ASN	771	20.278	1.636	-9.292	1.00 21.01
-	ATOM	791	ŏ	ASN	771	21.129	1.013	-8.648	1.00 20.52
	ATOM	792	N	GLU	772	19.258	1.043	-9.898	1.00 22.23
			CA	GLU	772	19.056	-0.393	-9.841	1.00 22.51
	ATOM	793			772	17.888		-10.711	1.00 23.17
	ATOM	794	CB	GLU				-10.099	1.00 24.81
55	ATOM	795	CG	GLU	772	16.562		-9.724	1.00 25.41
	ATOM	796	CD	GLU	772	15.761	-1.672	-9.724 -9.252	1.00 25.33
	MOTA	797		GLU	772	14.624	-1.488		
	ATOM	798		GLU	772	16.265	-2.803	-9.913	1.00 26.23
	ATOM	799	С	GLU	772	20.282	-1.102	-10.303	1.00 22.96
60	ATOM	800	0	GLU	772	20.631	-2.148	-9.785	1.00 23.89
	ATOM	801	N	TYR	773	20.961	-0.531	-11.276	1.00 22.51
	ATOM	802	CA	TYR	773	22.158	-1.164	-11.748	1.00 22.55
	-								

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	ATOM	803	CB 1	TYR	773	22.640	-0.492	-13.018	1.00 22	
	ATOM	804	CG 1	TYR	773	23.825		-13.593	1.00 22	
	ATOM	805	CD1	TYR	773	23.680		-14.304	1.00 23	
	ATOM	806		TYR	773	24.791	-3.041	-14.837		3.77
5	ATOM	807		TYR	773	25.095	-0.671	-13.418		2.59
•	ATOM	808	_	TYR	773	26.198	-1.309	-13.938	1.00 2	4.43
		809		TYR	773	26.047	-2.491	-14.643	1.00 2	4.22
	ATOM			TYR	773	27.172		-15.155	1.00 2	5.98
	ATOM	810		TYR	773	23.254		-10.680	1.00 2	3.17
40	ATOM	811		TYR	773	23.969		-10.523	1.00 2	4.08
10	ATOM	812	-			23.432	-0.044	-9.982	1.00 2	
	ATOM	813	-	ARG	774	24.427	0.047	-8.922		1.74
	MOTA	814	-	ARG	774		1.487	-8.494		2.23
	ATOM	815		ARG	774	24.623		-8.735	1.00 2	
	MOTA	816		ARG	774	26.026	1.952	-9.756		3.92
15	ATOM	817	_	ARG	774	26.073	3.066		1.00 2	
	ATOM	818	NE .	ARG	774	26.048	4.383	-9.146	1.00 2	
	ATOM	819		ARG	774	26.961	5.328	-9.365		
	ATOM	820	NH1	ARG	774	27.982	5.111	-10.171		5.01
	ATOM	821	NH2	ARG	774	26.837	6.509	-8.783	1.00 2	
20	ATOM	822	С	ARG	774	23.976	-0.796	-7.743		1.36
	ATOM	823	0	ARG	774	24.791	-1.386	-7.052	1.00 2	
	ATOM	824		MET	775	22.669	-0.854	-7.512	1.00 2	
	ATOM	825		MET	775	22.136	-1.681	-6.439	1.00 2	
	ATOM	826		MET	775	20.614	-1.582	-6.380	1.00 2	3.42
25		827		MET	775	20.121	-0.241	-5.955	1.00 2	3.46
25	ATOM	828		MET	775	18.333	-0.199	-5.865	1.00 2	6.50
	ATOM			MET	775	17.909	1.086	-7.064	1.00 2	7.26
	ATOM	829		MET	775	22.550	-3.136	-6.666	1.00 2	5.38
	ATOM	830			775	22.897	-3.832	-5.733		5.75
00	ATOM	831		MET	776	22.507	-3.593	-7.912	1.00 2	7.39
30	MOTA	832	N	HIS	776	22.891	-4.954	-8.262	1.00 2	
	ATOM	833	CA	HIS			-5.302	-9.684	1.00 2	
	MOTA	834	СВ	HIS	776	22.418		-10.067		10.57
	ATOM	835	CG	HIS	776	22.639		-9.864	1.00 3	
	ATOM	836	CD2		776	21.877	-7.843			80.81
35	MOTA	837	ND1		776	23.764		-10.739	1.00 2	
	MOTA	838		HIS	776	23.685		-10.932		
	ATOM	839	NE2	HIS	776	22.551		-10.411		9.53
	MOTA	840	С	HIS	776	24.403	-5.065	-8.178		9.55
	ATOM	841	0	HIS	776	24.923		-7.414		30.12
40	ATOM	842	N	LYS	777	25.109	-4.283	-8.989		31.13
	ATOM	843	CA	LYS	777	26.570	-4.290	-8.980		32.73
	ATOM	844	CB	LYS	777	27.130	-3.481	-10.161		31.29
	ATOM	845	CG	LYS	777	26.678	-3.948			30.55
	ATOM	846	CD	LYS	777	27.443		-12.003		29.83
45	ATOM	847	CE	LYS	777	28.928		-12.116		30.35
	ATOM	848	NZ	LYS	777	29.631	-5.860	-12.983	1.00 3	30.53
	ATOM	849	C	LYS	777	27.032	-3.655	-7.660	1.00 3	34.07
	ATOM	850	ŏ	LYS	777	27.382	-2.478	-7.611	1.00 3	
	ATOM	851	N	SER	778	26.995	-4.437	-6.596	1.00 3	33.74
50		852	CA	SER	778	27.387	-4.013		1.00 3	33.75
30	ATOM		СВ	SER	778	26.593	-2.789		1.00 3	
	ATOM	853			778	25.254	-3.122		1.00	
	ATOM	854	OG	SER	778	27.065	-5.204		1.00	
	ATOM	855	C	SER			-5.260		1.00	
	ATOM	856	0	SER	778	27.447	-6.149		1.00	
55	ATOM	857	N	ARG	779	26.344			1.00	
	ATOM	858	CA	ARG	779	25.926	-7.386		1.00	
	ATOM	859	CB	ARG	779	27.161	-8.256			
	ATOM	860	CG	ARG	779	28.065	-8.415		1.00	
	ATOM	861	CD	ARG	779	29.338	-9.182		1.00	
60	MOTA	862	NE	ARG	779	30.284	-9.129		1.00	
	ATOM	863	CZ	ARG	779	31.583	-9.401		1.00	
	ATOM	864	NH1	ARG	779	32.091	-9.753	-4.846	1.00	21.8/

	ATOM	865	NH2	ARG	779	32.398	-9.234	-7.050	1.00 26.64	
		866		ARG	779	25.128	-7.063	-3.084	1.00 30.75	
	ATOM		_	ARG	779	25.027	-7.880	-2.163	1.00 31.03	
	ATOM	867			780	24.521	-5.875	-3.097	1.00 29.82	
_	ATOM	868		MET			-5.381	-1.990	1.00 29.36	
5	ATOM	869		MET	780	23.721		-1.473	1.00 30.17	
	ATOM	870		MET	780	24.295	-4.068			
	ATOM	871	CG	MET	780	25.194	-4.191	-0.277		
	ATOM	872	SD	MET	780	25.835	-2.592	0.168	1.00 31.24	
	ATOM	873		MET	780	24.525	-1.995	1.114	1.00 31.13	
40				MET	780	22.262	-5.165	-2.331	1.00 29.21	
10	MOTA	874			780	21.542	-4.505	-1.566	1.00 29.43	
	ATOM	875		MET		21.831	-5.638	-3.497	1.00 28.58	
	ATOM	876		TYR	781			-3.897	1.00 28.82	
	ATOM	877	CA	TYR	781	20.433	-5.498		1.00 27.73	
	ATOM	878	CB	TYR	781	20.229	-5.985	-5.338		
15	ATOM	879	CG	TYR	781	18.896	-5.604	-5.964	1.00 26.05	
. •	ATOM	880	CD1		781	18.847	-4.861	-7.140	1.00 25.95	
		881	-	TYR	781	17.624	-4.510	-7.718	1.00 25.58	
	ATOM				781	17.686	-5.984	-5.382	1.00 25.08	
	ATOM	882	CD2			16.471	-5.643	-5.955	1.00 24.57	
	ATOM	883		TYR	781			-7.115	1.00 24.77	
20	MOTA	884	CZ	TYR	781	16.446	-4.904		1.00 24.60	
	ATOM	885	OH	TYR	781	15.238	-4.572	-7.668		
	ATOM	886	С	TYR	781	19.701	-6.393	-2.929	1.00 30.06	
	ATOM	887	Ö	TYR	781	19.984	-7.589	-2.856	1.00 32.08	
		888	N	SER	782	18.730	-5.821	-2.235	1.00 30.74	
25	ATOM			SER	782	17.935	-6.500	-1.198	1.00 31.41	
25	ATOM	889	CA		782	18.551	-7.836	-0.726	1.00 32.82	
	ATOM	890	СВ	SER			-8.438	0.308	1.00 35.91	
	ATOM	891	OG	SER	782	17.785			1.00 30.50	
	ATOM	892	С	SER	782	18.027	-5.483	-0.070		
	ATOM	893	0	SER	782	17.044	-4.807	0.230	1.00 30.31	
30	ATOM	894	N	GLN	783	19.242	-5.287	0.459	1.00 29.34	
00	ATOM	895	CA	GLN	783	19.455	-4.314	1.522	1.00 28.01	
	ATOM	896	CB	GLN	783	20.900	-4.309	2.020	1.00 28.50	í
		897	CG	GLN	783	21.327	-5.532	2.805	1.00 29.62	,
	ATOM				783	21.790	-6.634	1.900	1.00 32.01	
~=	ATOM	898	CD	GLN			-6.621	0.714	1.00 33.24	
35	ATOM	899			783	21.486		2.436	1.00 32.18	
	MOTA	900	NE2	GLN	783	22.547	-7.587			
	ATOM	901	С	GLN	783	19.089	-2.963	0.935	1.00 26.36	
	ATOM	902	0	GLN	783	18.342	-2.211	1.544	1.00 26.08	
	ATOM	903	N	CYS	784	19.538	-2.698	-0.290	1.00 25.49	
40	ATOM	904	CA	CYS	784	19.212	-1.439	-0.956	1.00 24.45	;
40				CYS	784	19.951	-1.312	-2.294	1.00 22.82	
	MOTA	905	CB		784	21.746	-0.989	-2.120	1.00 18.24	
	ATOM	906	SG	CYS			-1.290	-1.146	1.00 25.03	
	ATOM	907	С	CYS	784	17.698			1.00 25.67	
_	ATOM	908	0	CYS	784	17.155	-0.183	-1.044		
45	ATOM	909	N	VAL	785	17.003	-2.406	-1.360		
	ATOM	910	CA	VAL	785	15.538	-2.399	-1.547	1.00 25.02	
	ATOM	911	CB	VAL	785	14.987	-3.826	-1.903	1.00 25.94	
	ATOM	912		VAL	785	13.457	-3.901	-1.710	1.00 26.21	Ĺ
		913		VAL	785	15.349	-4.195	-3.324	1.00 26.31	Ĺ
EΩ	MOTA				785	14.864	-1.979	-0.257	1.00 23.91	
50	ATOM	914	C	VAL			-1.259	-0.260	1.00 24.19	
	ATOM	915	0	VAL	785	13.881			1.00 25.02	
	ATOM	916	N	ARG	786	15.402	-2.455	0.853		
	ATOM	917	CA	ARG	786	14.855	-2.158	2.165	1.00 25.39	
	ATOM	918	CB	ARG	786	15.468	-3.114	3.198	1.00 26.15	
55	ATOM	919	CG	ARG	786	15.392	-4.591	2.748	1.00 28.30)
55		920	CD	ARG	786	15.314	-5.583	3.900	1.00 29.76	
	ATOM					14.269	-5.206	4.851	1.00 32.39	
	ATOM	921	NE	ARG	786		-5.475	6.157	1.00 32.4	
	MOTA	922	CZ	ARG	786	14.292				
	ATOM	923		ARG	786	15.301	-6.153	6.701	1.00 32.09	
60	ATOM	924	NH2	ARG	786	13.326	-5.001	6.932	1.00 33.3	
	ATOM	925	С	ARG	786	15.083	-0.679	2.520	1.00 24.7	
	ATOM	926	0	ARG	786	14.180	-0.001	3.030	1.00 25.5	2
			-		-					

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	ATOM	927	N	MET	787 ·	16.246	-0.146	2.160	1.00 23.53
	ATOM	928	CA	MET	787	16.548	1.252	2.463	1.00 22.11
	ATOM	929	CB	MET	787	18.018	1.528	2.261	1.00 20.46
	ATOM	930	CG	MET	787	18.883	0.925	3.314	1.00 17.04
5	ATOM	931	SD	MET	787	20.578	0.861	2.788	1.00 20.46
•	ATOM	932	CE	MET	787	21.285	1.969	3.729	1.00 20.07
	ATOM	933	С	MET	787	15.736	2.173	1.588	1.00 23.39
	ATOM	934		MET	787	15.387	3.281	1.997	1.00 24.11
	ATOM	935		ARG	788	15.521	1.752	0.348	1.00 24.89
10	ATOM	936	CA	ARG	788	14.738	2.499	-0.625	1.00 26.29
	ATOM	937		ARG	788	14.833	1.790	-1.980	1.00 28.55
	ATOM	938	CG	ARG	788	14.166	2.474	-3.174	1.00 32.52
	ATOM	939	CD	ARG	788	14.217	1.541	-4.395	1.00 35.44
	ATOM	940	NE	ARG	788	13.426	1.996	-5.540	1.00 39.11
15	ATOM	941	CZ	ARG	788	13.899	2.177	-6.783	1.00 41.32
10	ATOM	942	NH1		788	15.182	1.960	-7.081	1.00 41.94
	ATOM	943	NH2		788	13.079	2.567	-7.754	1.00 41.48
	ATOM	944	С	ARG	788	13.312	2.475	-0.090	1.00 26.13
	ATOM	945	0	ARG	788	12.596	3.473	-0.146	1.00 26.50
20	ATOM	946	N	HIS	789	12.920	1.339	0.483	1.00 26.36
	ATOM	947	CA	HIS	789	11.587	1.173	1.052	1.00 26.76
	ATOM	948	СВ	HIS	789	11.377	-0.287	1.479	1.00 29.07
	ATOM	949	CG	HIS	789	9.970	-0.609	1.879	1.00 30.42
	ATOM	950	CD2	HIS	789	8.890	-0.944	1.137	1.00 31.35
25	ATOM	951	ND1	HIS	789	9.538	-0.567	3.188	1.00 32.05
	ATOM	952	CE1	HIS	789	8.249	-0.856	3.235	1.00 32.56
	ATOM	953		HIS	789	7.831	-1.087	2.001	1.00 32.55
	ATOM	954	С	HIS	789	11.369	2.133	2.231	1.00 26.08
	ATOM	955	0	HIS	789	10.275	2.671	2.394	1.00 25.72
30	ATOM	956	N	LEU	790	12.413	2.318	3.048	1.00 25.92
-	MOTA	957	CA	LEU	790	12.433	3.234	4.218	1.00 25.41
	ATOM	958	СВ	LEU	790	13.811	3.216	4.887	1.00 23.94
	ATOM	959	CG	LEU	790	14.039	3.400	6.383	1.00 23.32
	MOTA	960	CD1	LEU	790	15.444	3.930	6.570	1.00 22.41
35	ATOM	961	CD2	LEU	790	13.047	4.324	7.014	1.00 23.17
•	ATOM	962	С	LEU	790	12.218	4.654	3.720	1.00 25.47
	ATOM	963	0	LEU	790	11.359	5.380	4.216	1.00 25.06
	ATOM	964	N	SER	791	13.040	5.056	2.757	1.00 25.60
	ATOM	965	CA	SER	791	12.942	6.375	2.177	1.00 26.51
40	ATOM	966	CB	SER	791	13.851	6.446	0.973	1.00 28.35
	ATOM	967	OG	SER	791	14.936	5.559	1.179	1.00 32.32
	ATOM	968	С	SER	791	11.521	6.561	1.716	1.00 26.02
	ATOM	969	0	SER	7 9 1	10.950	7.632	1.885	1.00 26.00
	ATOM	970	N	GLN	792	10.964	5.505	1.122	1.00 26.31
45	ATOM	971	CA	GLN	792	9.600	5.526	0.610	1.00 26.32
	ATOM	972	CB	GLN	792	9.237	4.200	-0.112	1.00 28.65
	ATOM	973	CG	GLN	792	9.700	4.109	-1.603	1.00 30.43
	ATOM	974	CD	GLN	792	9.421	2.749	-2.277	1.00 31.95
	MOTA	975	OE1		792	8.479	2.607	-3.062	1.00 33.53
50	ATOM	976	NE2	GLN	792	10.273	1.764	-2.007	1.00 32.31
	ATOM	977	С	GLN	792	8.629	5.836	1.721	1.00 24.88
	ATOM	978	0	GLN	792	7.702	6.610	1.528	1.00 24.96
	ATOM	979	N	GLU	793	8.886	5.301	2.907	1.00 23.72
	MOTA	980	CA	GLU	793	8.014	5.550	4.051	1.00 22.89
55	ATOM	981	CB	GLU	793	8.460	4.728	5.273	1.00 23.65
	ATOM	982	CG	GLU	793	8.555	3.199	5.055	1.00 25.18
	ATOM	983	CD	GLU	793	7.383	2.406	5.651	1.00 27.08
	ATOM	984		GLU	793	6.207	2.735	5.351	1.00 25.97
	MOTA	985	OE2	GLU	793	7.648	1.450	6.433	1.00 28.69
60	ATOM	986	С	GLU	793	7.949	7.041	4.400	1.00 21.58
	MOTA	987	0	GLU	793	6.903	7.530	4.764	1.00 21.52
	MOTA	988	N	PHE	794	9.042	7.784	4.274	1.00 21.26

	ATOM	989	CA PHE	794		8.999	9.208	4.598	1.00 20.65 1.00 19.81
	ATOM	990	CB PHE	794		0.334	9.890	4.323 5.304	1.00 19.96
	ATOM	991	CG PHE	794		1.413	9.541	6.662	1.00 20.01
	ATOM	992	CD1 PHE	794		1.226	9.728	4.878	1.00 20.01
5	ATOM	993	CD2 PHE	794		2.599	8.974	7.566	1.00 19.86
_	ATOM	994	CE1 PHE	794		2.206	9.347	5.787	1.00 18.95
	MOTA	995	CE2 PHE	794		3.570	8.593		1.00 10.33
	ATOM	996	CZ PHE	794	1	3.374	8.777	7.118	1.00 19.37
	MOTA	997	C PHE	794		7.929	9.863	3.759	1.00 22.20
10	ATOM	998	O PHE	794		7.387	10.906	4.138	
	ATOM	999	N GLY	795		7.688	9.270	2.585	
	ATOM	1000	CA GLY	795		6.676	9.750	1.662	
	ATOM	1001	C GLY	795		5.309	9.232	2.037	1.00 26.19
	ATOM	1002	O GLY	795		4.414	10.002	2.345	1.00 27.46
15	ATOM	1003	N TRP	796		5.181	7.912	2.081	1.00 27.45
, 0	ATOM	1004	CA TRP	796		3.931	7.239	2.428	1.00 28.24
	ATOM	1005	CB TRP	796		4.135	5.697	2.542	1.00 27.71
	ATOM	1006	CG TRP	796		4.478	4.998	1.187	1.00 27.50
	ATOM	1007	CD2 TRP	796		5.208	3.763	0.985	1.00 26.97
20	MOTA	1008	CE2 TRP	796		5.312	3.556	-0.417	1.00 26.72
20	ATOM	1009	CE3 TRP	796		5.777	2.816	1.845	1.00 25.52
	ATOM	1010	CD1 TRP	796		4.177	5.460	-0.079	1.00 27.17
	ATOM	1011	NE1 TRP	796		4.676	4.601	-1.035	1.00 27.59
	ATOM	1012	CZ2 TRP	796		5.967	2.448	-0.970	1.00 25.70
25		1012	CZ3 TRP	796		6.427	1.714	1.290	1.00 25.51
25	ATOM	1013	CH2 TRP	796		6.514	1.543	-0.106	1.00 25.42
	ATOM ATOM	1015	C TRP			3.345	7.826	3.706	1.00 29.13
		1015	O TRP			2.132	8.026	3.801	1.00 29.87
	ATOM	1017	N LEU			4.223	8.212	4.632	1.00 29.96
30	ATOM	1017	CA LEU			3.816	8.768	5.923	1.00 29.80
30	ATOM	1019	CB LEU			4.692	8.223	7.061	1.00 28.43
	ATOM	1019	CG LEU			4.552	6.736	7.383	1.00 27.68
	ATOM	1020	CD1 LEU			5.709	6.269	8.228	1.00 27.20
	ATOM	1021	CD2 LEU			3.216	6.470	8.058	1.00 26.62
25	ATOM	1022	C LEU			3.864	10.260	5.991	1.00 30.39
35	ATOM					3.447	10.827	6.983	1.00 32.25
	ATOM	1024				4.415	10.908	4.978	1.00 31.03
	ATOM	1025	-			4.518	12.360	5.005	1.00 30.93
	ATOM	1026				3.117	13.030	4.964	1.00 31.58
40	ATOM	1027	CB GLN			2.253	12.757	3.701	1.00 32.86
40	ATOM	1028	CG GLN			0.944	13.580	3.633	1.00 32.89
	MOTA	1029				0.342	13.933	4.648	1.00 33.16
	ATOM	1030				0.521	13.892	2.421	1.00 33.46
	MOTA	1031	NE2 GLN			5.267	12.764	6.294	1.00 30.02
45	MOTA	1032	C GLN			4.716	13.460	7.147	1.00 30.51
45	MOTA	1033	O GLN			6.497	12.283	6.462	1.00 28.60
	MOTA	1034	N ILE				12.634	7.648	1.00 27.76
	MOTA	1035	CA ILE			7.277 8.546		7.829	1.00 26.34
	MOTA	1036	CB ILE			9.382	12.246	9.007	1.00 25.78
	ATOM	1037	CG2 ILE					8.046	1.00 25.57
50	ATOM	1038	CG1 ILE			8.168		9.211	1.00 25.52
	ATOM	1039	CD1 ILE			7.271		7.552	1.00 28.59
	ATOM	1040	C ILE			7.729		6.496	1.00 29.81
	MOTA	1041	O ILE			8.181			1.00 28.71
	ATOM	1042	N THE			7.610		8.678	1.00 28.68
55	MOTA	1043	CA THE			7.967		8.874	1.00 28.00
	ATOM	1044	CB THE			7.202		10.142	
	MOTA	1045	OG1 THE			5.839		9.815	1.00 31.49
	ATOM	1046	CG2 TH			7.824		10.746	1.00 29.69
	MOTA	1047	C THI			9.475		9.116	1.00 28.69
60	ATOM	1048	O THI			10.069		9.796	1.00 29.15
	ATOM	1049				10.116		8.565	1.00 28.18
	ATOM	1050	CD PRO	801		9.618	18.376	7.569	1.00 27.62

								0 700	1.00 27.74
	MOTA	1051	CA	PRO	801	11.555	17.600	8.780 8.178	1.00 27.74
	MOTA	1052	CB	PRO	801	11.797	18.983	7.002	1.00 26.58
	ATOM	1053	CG	PRO	801	10.908	18.956	10.271	1.00 27.38
	MOTA	1054	С	PRO	801	11.907	17.570 17.101	10.666	1.00 27.75
5	MOTA	1055	0	PRO	801	12.981	18.045	11.095	1.00 27.01
	MOTA	1056	N	GLN	802	10.982 11.189	18.079	12.542	1.00 26.73
	ATOM	1057	CA	GLN	802	10.316	19.162	13.192	1.00 28.09
	ATOM	1058	CB	GLN	802	10.510	20.596	12.692	1.00 29.79
	ATOM	1059	CG	GLN	802	9.997	20.900	11.303	1.00 30.48
10	ATOM	1060	CD	GLN	802 802	8.948	20.381	10.918	1.00 30.36
	MOTA	1061	OE1		802 802	10.660	21.782	10.571	1.00 30.57
	ATOM	1062	NE2		802	10.968	16.715	13.219	1.00 24.81
	MOTA	1063	C	GLN GLN	802	11.599	16.415	14.222	1.00 24.41
4.5	ATOM	1064	0		803	10.064	15.904	12.669	1.00 23.52
15	ATOM	1065	N	GLU GLU	803	9.797	14.558	13.196	1.00 21.64
	ATOM	1066	CA	GLU	803	8.632	13.897	12.459	1.00 20.29
	ATOM	1067 1068	CB CG	GLU	803	7.277	14.434	12.848	1.00 18.44
	ATOM	1069	CD	GLU	803	6.147	13.786	12.119	1.00 17.84
20	ATOM	1070		GLU	803	6.308	13.392	10.958	1.00 18.19
20	MOTA	1070		GLU	803	5.065	13.680	12.704	1.00 19.88
	ATOM	1071	C	GLU	803	11.067	13.784	12.923	1.00 21.04
	ATOM ATOM	1072	Ö	GLU	803	11.537	13.042	13.777	1.00 20.89
	ATOM	1074	N	PHE	804	11.612	14.001	11.722	1.00 19.87
25	ATOM	1075	CA	PHE	804	12.863	13.418	11.254	1.00 19.24
20	ATOM	1076	СВ	PHE	804	13.144	13.867	9.822	1.00 17.23
	ATOM	1077	CG	PHE	804	14.557	13.645	9.384	1.00 14.85
	ATOM	1078		PHE	804	15.012	12.380	9.095	1.00 13.76
	ATOM	1079	CD2	PHE	804	15.440	14.706	9.301	1.00 13.69
30	ATOM	1080	CE1	PHE	804	16.335	12.160	8.729	1.00 13.79
••	ATOM	1081	CE2	PHE	804	16.765	14.496	8.936	1.00 13.36
	MOTA	1082	CZ	PHE	804	17.214	13.217	8.647	1.00 12.84
	ATOM	1083	С	PHE	804	14.034	13.802	12.157	1.00 20.31
	ATOM	1084	0	PHE	804	14.807	12.939	12.564	1.00 21.30
35	ATOM	1085	N	LEU	805	14.187	15.086	12.463	1.00 20.30 1.00 20.09
	MOTA	1086	CA	LEU	805	15.271	15.503	13.339 13.582	1.00 20.03
	ATOM	1087	CB	LEU	805	15.250	17.008 17.834	12.330	1.00 19.30
	MOTA	1088	CG	LEU	805	15.552	19.281	12.707	1.00 19.84
40	MOTA	1089		LEU	805	15.704	17.343	11.670	1.00 19.41
40	MOTA	1090		LEU	805	16.816 15.172	14.767	14.651	1.00 19.83
	ATOM	1091	С	LEU	805 805	16.142	14.205	15.106	1.00 20.77
	MOTA	1092	0	LEU	806	13.980	14.719	15.223	1.00 20.17
	ATOM	1093 1094	N CA	CYS CYS	806	13.765	14.026	16.494	1.00 21.27
45	ATOM ATOM	1094	CB	CYS	806	12.372	14.332	17.078	1.00 22.13
45		1095	SG	CYS	806	12.142	16.017	17.706	1.00 27.50
	ATOM ATOM	1097	C	CYS	806	13.938	12.515	16.378	1.00 20.36
	ATOM	1098	Ö	CYS	806	14.575	11.904	17.241	1.00 20.30
	ATOM	1099	N	MET	807	13.348	11.903	15.350	1.00 19.67
50	ATOM	1100	CA	MET	807	13.491	10.458	15.160	1.00 18.10
00	ATOM	1101	CB	MET	807	12.668	9.944	13.989	1.00 17.25
	ATOM	1102	CG	MET	807	11.195	9.877	14.279	1.00 16.70
	ATOM	1103	SD	MET	807	10.377	9.142	12.911	1.00 19.42
	ATOM	1104	CE	MET	807	10.144	10.560	11.908	1.00 16.21
55	ATOM	1105	С	MET	807	14.947	10.062	14.979	1.00 17.75
	ATOM	1106	0	MET	807	15.371	9.038	15.490	1.00 18.41
	ATOM	1107	N	LYS	808	15.712	10.871	14.257	1.00 17.31
	ATOM	1108	CA	LYS	808	17.116	10.592	14.054	1.00 16.27
	ATOM	1109	СВ	LYS	808	17.729	11.514	12.994	1.00 15.01
60	MOTA	1110	CG	LYS	808	19.171	11.154	12.733	1.00 14.63
	ATOM	1111	CD	LYS	808	19.679	11.569	11.371	1.00 15.42
	MOTA	1112	ÇE	LYS	808	19.422	13.053	11.092	1.00 15.64

1.00 14.15 20.232 13.940 11.928 808 LYS 1113 ΝZ **ATOM** 1.00 16.89 15.376 10.726 17.857 808 LYS MOTA 1114 C 1.00 16.07 18.731 9.908 15.677 LYS 808 0 ATOM 1115 1.00 16.53 16.166 17.522 11.747 809 1116 N ALA MOTA 1.00 17.68 17.461 18.175 11.931 ALA 809 CA 1117 **ATOM** 18.155 1.00 16.91 17.628 13.139 809 1118 CB ALA MOTA 1.00 19.03 18.348 17.989 10.691 809 ALA С ATOM 1119 1.00 20.50 10.207 18.996 18.932 809 ALA 1120 0 ATOM 1.00 19.36 18.392 10.184 16.766 810 LEU 1121 N ATOM 1.00 18.99 16.459 9.011 19.186 810 LEU 10 1122 CA ATOM 1.00 19.09 14.966 8.811 19.263 810 LEU CB 1.00 20.20 1.00 21.29 1.00 18.75 1.00 18.88 1123 **ATOM** 20.651 9.020 810 14.406 CG LEU MOTA 1124 12.954 20.594 8.606 810 CD1 LEU ATOM 1125 21.674 15.176 8.199 CD2 LEU 810 1126 ATOM 7.716 18.722 17.116 810 С LEU 15 ATOM 1127 1.00 20.78 6.780 19.509 17.213 810 LEU 0 1128 ATOM 1.00 17.63 17.456 17.537 7.636 1129 N LEU 811 MOTA 1.00 15.58 16.959 811 6.447 18.215 CA LEU A'TOM 1130 1.00 14.70 18.346 6.456 15.438 CB LEU 811 MOTA 1131 1.00 14.14 1.00 13.66 14.574 6.107 811 17.148 LEU CG 20 ATOM 1132 13.164 14.746 6.408 17.511 CD1 LEU 811 MOTA 1133 1.00 13.62 16.744 4.632 CD2 LEU 811 MOTA 1134 17.582 1.00 15.42 19.598 6.328 811 LEU 1135 С ATOM 1.00 17.27 20.189 5.252 17.554 811 LEU 0 MOTA 1136 1.00 13.97 18.084 7.429 20.153 25 N LEU 812 1137 ATOM 7.373 18.734 1.00 12.94 21.455 812 LEU MOTA 1138 CA 1.00 12.69 18.937 22.004 8.790 LEU 812 CB **ATOM** 1139 1.00 12.03 19.670 8.893 23.342 812 LEU 1140 CG MOTA 1.00 12.16 18.802 24.488 8.422 CD1 LEU 812 MOTA 1141 1.00 13.12 20.037 23.559 10.325 CD2 LEU 812 30 1142 1.00 12.97 1.00 12.99 1.00 13.55 MOTA 6.658 20.098 21.330 812 LEU **ATOM** 1143 С 6.118 20.629 6.681 20.662 6.064 21.950 LEU 812 22.282 0 **ATOM** 1144 20.136 813 MOTA 1145 N PHE 1.00 14.19 CA PHE 813 19.859 1146 ATOM 1.00 15.20 7.088 22.821 19.137 PHE 813 35 ÇВ 1147 **ATOM** 1.00 16.11 8.435 22.841 19.818 813 CG PHE ATOM 1148 8.640 23.624 9.472 22.036 1.00 15.97 20.946 CD1 PHE 813 MOTA 1149 1.00 16.07 19.349 CD2 PHE 813 1150 ATOM 1.00 18.37 9.845 23.615 21.604 813 CE1 PHE 1151 MOTA 1.00 17.54 10.687 22.014 19.991 40 CE2 PHE 813 1152 MOTA 1.00 17.99 10.883 22.801 21.126 813 CZ PHE MOTA 1153 1.00 14.71 18.971 4.856 21.753 813 1154 PHE ATOM C 1.00 14.79 4.618 22.530 18.058 PHE 813 1155 0 **ATOM** 1.00 16.09 20.709 4.082 19.255 814 ATOM 1156 N SER 18.453 20.369 1.00 15.96 2.917 CA SER 814 45 1157 ATOM 1.00 15.79 19.062 17.697 3.172 MOTA 814 1158 CB SER 19.274 1.00 15.51 4.087 814 16.640 1159 OG SER MOTA 1.00 16.05 19.169 1.581 20.294 1160 C SER 814 MOTA 1.00 17.02 19.779 0.620 814 18.610 1161 0 SER **ATOM** 20.779 1.00 16.07 20.395 1.498 815 50 1162 N ILE MOTA 1.00 17.04 20.747 21.099 0.226 1163 CA ILE 815 **ATOM** 1.00 16.75 1.00 17.43 19.325 -0.086 CB ILE 21.620 815 MOTA 1164 1.113 22.222 18.706 CG2 ILE 815 1165 MOTA 1.00 17.01 -1.245 -1.753 19.341 22.600 1166 CG1 ILE 815 MOTA 17.953 1.00 17.98 22.915 1167 CD1 ILE 815 55 MOTA 1.00 17.86 0.187 21.826 22.172 815 1168 C ILE MOTA 1.00 18.49 1169 0 0.981 21.802 23.111 815 ILE MOTA -0.700 22.809 1.00 18.25 21.994 ILE 816 1170 N MOTA 1.00 18.91 23.947 22.913 -0.804 1171 CA ILE 816 ATOM 25.178 1.00 19.07 816 22.298 816 22.175 816 20.939 CB ILE -0.099 60 1172 MOTA 1.378 24.921 1.00 17.73 1173 CG2 ILE MOTA 1.00 18.04 25.537 -0.692 1174 CG1 ILE **ATOM**

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									1.00 17.73
	MOTA	1175	CD1	ILE	816	20.516	-0.346	26.933	1.00 17.73
	ATOM	1176	C	ILE	816	23.302	-2.226	24.385	_
	ATOM	1177	0	ILE	816	22.615	-3.184	24.040	1.00 20.43 1.00 20.85
	ATOM	1178	N	PRO	817	24.392	-2.385	25.180	
5	ATOM	1179	CD	PRO	817	25.303	-1.373	25.730	1.00 21.00
_	ATOM	1180	CA	PRO	817	24.805	-3.720	25.631	1.00 22.05
	MOTA	1181	CB	PRO	817	26.016	-3.444	26.523	1.00 21.47
	ATOM	1182	CG	PRO	817	26.554	-2.197	26.001	1.00 21.80
	ATOM	1183	С	PRO	817	23.706	-4.320	26.458	1.00 22.95
10	ATOM	1184	0	PRO	817	22.988	-3.594	27.151	1.00 23.12
	ATOM	1185	N	VAL	818	23.585	-5.640	26.418	1.00 24.79
	ATOM	1186	CA	VAL	818	22.544	-6.316	27.195	1.00 26.35
	ATOM	1187	CB	VAL	818	22.513	-7.860	26.916	1.00 27.19 1.00 27.82
	ATOM	1188	CG1		818	23.864	-8.515	27.282	
15	ATOM	1189	CG2	VAL	818	21.362	-8.524	27.676	1.00 27.72
-	ATOM	1190	С	VAL	818	22.742	-6.047	28.692	1.00 26.79
	ATOM	1191	0	VAL	818	21.777	-5.849	29.421	1.00 26.79
	ATOM	1192	N	ASP	819	23.992	-5.963	29.136	1.00 27.83
	ATOM	1193	CA	ASP	819	24.240	-5.732	30.550	1.00 29.78
20	ATOM	1194	CB	ASP	819	25.406	-6.593	31.063	1.00 32.59
	ATOM	1195	CG	ASP	819	26.747	-5.908	30.959	1.00 35.35
	ATOM	1196	OD1	ASP	819	27.117	-5.518	29.825	1.00 38.62
	ATOM	1197	OD2	ASP	819	27.431	-5.776	32.011	1.00 36.18 1.00 29.73
	ATOM	1198	С	ASP	819	24.377	-4.266	30.937	
25	ATOM	1199	0	ASP	819	24.899	-3.930	32.007	1.00 30.00 1.00 29.43
	ATOM	1200	N	GLY	820	23.839	-3.403	30.085	
	ATOM	1201	CA	GLY	820	23.878	-1.974	30.342	1.00 28.69 1.00 27.42
	ATOM	1202	С	GLY	820	25.216	-1.317	30.125 29.938	1.00 27.42
	ATOM	1203	0	GLY	820	26.221	-1.982		1.00 28.29
30	MOTA	1204	N	LEU	821	25.208	0.010	30.135	1.00 28.64
	ATOM	1205	CA	LEU	821	26.410	0.831	29.947 29.195	1.00 28.29
	ATOM	1206	CB	LEU	821	26.023	2.110	27.991	1.00 28.23
	ATOM	1207	CG	LEU	821	25.083	1.940	28.031	1.00 27.27
	ATOM	1208	CD1		821	24.046	3.022	26.653	1.00 27.18
35	ATOM	1209	CD2		821	25.831	1.953	31.349	1.00 27.10
	ATOM	1210	C	LEU	821	26.948	0.747	32.342	1.00 28.84
	MOTA	1211	0	LEU	821	26.341 28.060	1.897	31.441	1.00 28.49
	ATOM	1212	N	LYS	822	28.642	2.268	32.741	1.00 29.80
40	MOTA	1213	CA	LYS	822	29.865	3.169	32.576	1.00 30.45
40	ATOM	1214	CB	LYS	822 822	30.924	2.626	31.666	1.00 32.84
	ATOM	1215	CD	LYS LYS	822	31.517	1.345	32.194	1.00 35.27
	ATOM	1216 1217	CE	LYS	822	32.433	0.688	31.161	1.00 36.20
	ATOM	1217	NZ	LYS	822	33.498	1.623	30.710	1.00 37.22
45	ATOM ATOM	1218	C	LYS	822	27.621	3.016	33.587	1.00 30.25
45	ATOM	1220	0	LYS	822	27.353	2.655	34.731	1.00 31.02
	ATOM	1221	N	ASN	823	27.065	4.080	33.029	1.00 29.98
	ATOM	1222	CA	ASN	823	26.070	4.852	33.735	1.00 29.55
	ATOM	1223	CB	ASN	823	26.458	6.323	33.774	1.00 31.17
50	ATOM	1224	CG	ASN	823	27.832	6.544	34.350	1.00 32.55
00	ATOM	1225		ASN	823	28.787	5.856	33.985	1.00 33.56
	ATOM	1226		ASN	823	27.952	7.520	35.246	1.00 34.42
	ATOM	1227	С	ASN	823	24.807	4.665	32.943	1.00 28.73
	ATOM	1228	ō	ASN	823	24.476	5.473	32.091	1.00 29.00
55	MOTA	1229	N	GLN	824	24.127	3.562	33.199	1.00 27.99
	ATOM	1230	CA	GLN	824	22.893	3.227	32.514	1.00 27.77
	ATOM	1231	CB	GLN	824	22.590	1.731	32.738	1.00 28.13
	ATOM	1232	CG	GLN	824	21.343	1.158	32.077	1.00 28.93
	ATOM	1233	CD	GLN	824	21.331	1.302	30.551	1.00 30.20
60	ATOM	1234		GLN	824	22.300	0.976	29.855	1.00 30.02
	ATOM	1235		GLN	824	20.211	1.775	30.028	1.00 29.72
	ATOM	1236	С	GLN	824	21.723	4.115	32.960	1.00 27.61

						00			
	MOTA	1237	0	GLN	824	20.747	4.275	32.226	1.00 27.58
	ATOM	1238	N	LYS	825	21.833	4.752	34.122	1.00 27.13
	ATOM	1239	CA	LYS	825	20.742	5.590	34.595	1.00 26.37
	ATOM	1240	CB	LYS	825	20.815	5.802	36.113	1.00 28.08
5	ATOM	1241	CG	LYS	825	19.430	5.823	36.792	1.00 31.02
•	ATOM	1242	CD	LYS	825	19.493	5.693	38.335	1.00 33.41
	ATOM	1243	CE	LYS	825	18.086	5.725	39.002	1.00 34.57
	ATOM	1244	NZ	LYS	825	17.196	4.516	38.739	1.00 35.55
	ATOM	1245	С	LYS	825	20.679	6.917	33.876	1.00 24.74
10	ATOM	1246	0	LYS	825	19.625	7.518	33.799	1.00 25.30
	ATOM	1247	N	PHE	826	21.794	7.375	33.330	1.00 24.06
	ATOM	1248	CA	PHE	826	21.830	8.646	32.597	1.00 23.44 1.00 25.61
	ATOM	1249	CB	PHE	826	23.247	9.191	32.573	1.00 28.86
	ATOM	1250	CG	PHE	826	23.768	9.527	33.930	1.00 29.49
15	MOTA	1251		PHE	826	22.916	10.067	34.890	1.00 29.49
	ATOM	1252		PHE	826	25.091	9.284	34.268	1.00 29.57
	MOTA	1253		PHE	826	23.373	10.356	36.156	1.00 29.80
	ATOM	1254		PHE	826	25.551	9.571	35.533	1.00 29.00
	MOTA	1255	CZ	PHE	826	24.688	10.108	36.479 31.178	1.00 30.14
20	MOTA	1256	С	PHE	826	21.344	8.463	30.568	1.00 21.42
	ATOM	1257	0	PHE	826	20.808	9.380 7.277	30.636	1.00 20.42
	ATOM	1258	N	PHE	827	21.581	6.937	29.299	1.00 18.74
	ATOM	1259	CA	PHE	827	21.145	5.644	28.857	1.00 17.62
	MOTA	1260	CB	PHE	827	21.814 21.238	5.083	27.610	1.00 16.77
25	MOTA	1261	CG	PHE	827	21.780	5.412	26.380	1.00 16.74
	MOTA	1262		PHE	827	20.123	4.261	27.656	1.00 16.46
	ATOM	1263		PHE	827	21.225	4.939	25.212	1.00 16.50
	MOTA	1264	CE1		827 827	19.555	3.782	26.491	1.00 17.41
20	MOTA	1265	CE2		827	20.105	4.120	25.266	1.00 16.35
30	ATOM	1266	CZ	PHE	827	19.627	6.778	29.277	1.00 19.19
	ATOM	1267	C	PHE	827	18.962	7.183	28.331	1.00 18.79
	MOTA	1268	0 N	ASP	828	19.079	6.150	30.312	1.00 20.42
	ATOM	1269 1270	CA	ASP	828	17.638	5.943	30.421	1.00 21.69
35	ATOM ATOM	1271	CB	ASP	828	17.325	5.045	31.633	1.00 23.37
33	ATOM	1272	CG	ASP	828	17.885	3.627	31.487	1.00 24.46
	ATOM	1273		ASP	828	17.900	3.095	30.365	1.00 26.34
	MOTA	1274		ASP	828	18.296	3.023	32.501	1.00 26.77
	ATOM	1275	Č	ASP	828	16.931	7.287	30.572	1.00 21.41
40	ATOM	1276	Ö	ASP	828	15.835	7.487	30.070	1.00 21.87
	ATOM	1277	N	GLU	829	17.552	8.187	31.313	1.00 22.11
	ATOM	1278	CA	GLU	829	17.005	9.510	31.533	1.00 23.92
	ATOM	1279	CB	GLU	829	17.910	10.309	32.499	1.00 27.77
	ATOM	1280	CG	GLU	829	18.168	11.823	32.130	1.00 32.20
45	ATOM	1281	CD	GLU	829	19.650	12.266	32.334	1.00 35.29
	ATOM	1282	OE1	GLU	829	20.005	12.655	33.482	1.00 37.06
	ATOM	1283	OE2	GLU	829	20.463	12.217	31.360	1.00 34.89
	MOTA	1284	С	GLU	829	17.011	10.166	30.174	1.00 22.78
	ATOM	1285	0	GLU	829	15.963	10.539	29.656	1.00 22.06
50	MOTA	1286	N	LEU	830	18.201	10.200	29.575	1.00 22.21 1.00 22.14
	MOTA	1287	CA	LEU	830	18.437	10.812	28.272	1.00 22.14
	MOTA	1288	CB	LEU	830	19.885	10.575	27.852	1.00 21.24
	MOTA	1289	CG	LEU	830	20.415	11.572	26.833	1.00 21.40
	ATOM	1290		LEU	830	20.037	13.004	27.215	1.00 21.40
55	ATOM	1291		LEU	830	21.895	11.429	26.752 27.191	1.00 22.34
	ATOM	1292	C	LEU	830	17.499	10.318		1.00 23.35
	MOTA	1293	0	LEU	830	16.874	9.002	26.481 27.079	1.00 23.33
	ATOM	1294	N	ARG	831	17.400	8.352	26.097	1.00 22.23
60	MOTA	1295	CA	ARG	831	16.559 16.780	6.849	26.186	1.00 22.50
60	ATOM	1296	CB	ARG	831 831	15.760	6.087	25.219	1.00 22.59
	ATOM	1297	CG	ARG	831	16.130	4.600	25.375	1.00 23.29
	ATOM	1298	CD	ARG	0.01	10.130	1.500		

							2 272	24 074	1.00 25.50
	ATOM	1299		ARG	831	15.921	3.972	24.074	1.00 24.36
	MOTA	1300	_	ARG	831	14.738	3.800	23.491	1.00 24.30
	MOTA	1301	NH1	ARG	831	13.632	4.173	24.096	1.00 24.31
	ATOM	1302	NH2	ARG.	831	14.676	3.366	22.250	1.00 24.31
5	ATOM	1303	С	ARG	831	 15.086	8.667	26.302	
	ATOM	1304	0	ARG	831	14.354	8.964	25.351	1.00 22.89 1.00 22.76
	ATOM	1305	N	MET	832	14.655	8.593	27.550	
	ATOM	1306	CA	MET	832	13.276	8.859	27.923	1.00 22.91
	ATOM	1307	СВ	MET	832	13.126	8.762	29.429	1.00 23.88
10	ATOM	1308	CG	MET	832	11.739	9.050	29.870	1.00 24.65
10	ATOM	1309	SD	MET	832	11.693	9.332	31.596	1.00 29.43
	ATOM	1310	CE	MET	832	10.059	10.026	31.651	1.00 29.67
	ATOM	1311	c	MET	832	12.879	10.262	27.513	1.00 23.47
	ATOM	1312	ŏ	MET	832	11.740	10.512	27.097	1.00 23.79
15		1313	N	ASN	833	13.782	.11.198	27.768	1.00 23.63
13	ATOM	1314	CA	ASN	833	13.562	12.599	27.423	1.00 23.80
	ATOM	1315	CB	ASN	833	14.676	13.482	28.013	1.00 23.64
	ATOM	1316	CG	ASN	833	14.532	13.679	29.544	1.00 23.87
	MOTA	1317	OD1		833	15.519	13.864	30.270	1.00 23.24
20	MOTA	1317	ND2		833	13.293	13.628	30.030	1.00 24.57
20	MOTA		C	ASN	833	13.403	12.761	25.905	1.00 23.94
	ATOM	1319	Ö	ASN	833	12.463	13.397	25.445	1.00 24.48
	ATOM	1320	N	TYR	834	14.240	12.093	25.123	1.00 23.69
	ATOM	1321	CA	TYR	834	14.121	12.165	23.673	1.00 24.70
25	ATOM	1322	CB	TYR	834	15.340	11.532	23.007	1.00 25.39
25	ATOM	1323	CG	TYR	834	16.491	12.489	22.872	1.00 25.49
	ATOM	1324		TYR	834	16.802	13.051	21.635	1.00 26.75
	MOTA	1325	CD1	TYR	834	17.828	13.975	21.502	1.00 27.67
	ATOM	1326			834	17.239	12.869	23.986	1.00 26.00
00	ATOM	1327	CD2		834	18.268	13.791	23.873	1.00 26.69
30	ATOM	1328	CE2		834	18.558	14.341	22.624	1.00 28.15
	ATOM	1329	CZ	TYR	834	19.571	15.263	22.497	1.00 28.86
	MOTA	1330	OH	TYR	834	12.809	11.574	23.128	1.00 24.43
	MOTA	1331	C	TYR	834	12.297	12.006	22.082	1.00 24.26
~=	ATOM	1332	0	TYR	835	12.260	10.599	23.843	1.00 24.33
35	MOTA	1333	N	ILE	835	11.004	9.991	23.450	1.00 23.48
	MOTA	1334	CA	ILE	835	10.724	8.727	24.235	1.00 21.33
	MOTA	1335	CB	ILE	835	9.297	8.308	24.054	1.00 19.87
	ATOM	1336	CG2		835	11.657	7.624	23.756	1.00 20.40
40	MOTA	1337	CG1		835	11.684	6.400	24.655	1.00 21.05
40	MOTA	1338	CD1		835	9.893	10.977	23.687	1.00 25.05
	MOTA	1339	C	ILE	835	8.972	11.061	22.889	1.00 26.24
	ATOM	1340	0	ILE	836	9.998	11.738	24.771	1.00 26.82
	ATOM	1341	N	LYS	836	9.006	12.747	25.137	1.00 28.28
45	ATOM	1342	CA	LYS	836	9.245	13.281	26.556	1.00 29.95
45	ATOM	1343	CB	LYS	836	9.115	12.252	27.712	1.00 32.62
	ATOM	1344	CG	LYS	836	7.690	11.672	27.901	1.00 33.04
	ATOM	1345	CD	LYS		7.575	10.238	27.344	1.00 34.32
	ATOM	1346	CÉ	LYS	836	8.559	9.259	27.942	1.00 32.90
	MOTA	1347	NZ	LYS	836 836	9.030	13.919	24.164	1.00 28.39
50	ATOM	1348	C	LYS	836	7.997	14.545	23.946	1.00 29.09
	MOTA	1349	0	LYS		10.194	14.249	23.606	1.00 28.61
	MOTA	1350	N	GLU	837 837	10.134	15.351	22.643	1.00 28.70
	MOTA	1351	CA	GLU		11.715	15.824	22.439	1.00 29.34
	MOTA	1352	CB	GLU	837	12.305	16.584	23.627	1.00 32.13
55	ATOM	1353	CG	GLU	837		17.887	23.027	1.00 34.30
	ATOM	1354	CD	GLU	837	11.553	18.303	25.157	1.00 34.18
	MOTA	1355		GLU	837	11.612	18.503		
	MOTA	1356		GLU	837	10.925	14.907		1.00 28.92
	MOTA	1357		GLU	837	9.666	15.699		
60	MOTA	1358	0	GLU	837	9.041	13.631		1.00 29.32
	ATOM	1359		LEU	838	9.826 9.250	13.031		1.00 30.90
	ATOM	1360	CA	LEU	838	3.430	17.032	23.774	2.00 00.00

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								10 502	1.00 30.44
	ATOM	1361		LEU	838	9.614	11.622	19.592	1.00 30.56
	ATOM	1362		LEU	838	8.810	10.983	18.460	1.00 30.30
	MOTA	1363	CD1		838	9.077	11.728	17.151	1.00 30.31
	ATOM	1364	CD2	LEU	838	9.166	9.533	18.330	1.00 30.00
5	MOTA	1365	_	LEU	838	7.740	13.211	19.884	
	MOTA	1366	0	LEU	838	7.076	13.706	18.983	
	ATOM	1367	N .	ASP	839	7.186	12.724	20.979	1.00 34.82
	ATOM	1368	CA	ASP	839	5.755	12.823	21.162	1.00 37.61
	ATOM	1369	CB .	ASP	839	5.331	12.117	22.449	1.00 39.22
10	ATOM	1370	CG .	ASP	839	3.816	11.952	22.557	1.00 41.54
. •	ATOM	1371	OD1	ASP	839	3.249	12.375	23.592	1.00 43.10
	ATOM	1372	OD2		839	3.192	11.400	21.613	1.00 42.60
	ATOM	1373		ASP	839	5.338	14.293	21.187	1.00 38.83
	ATOM	1374		ASP	839	4.285	14.645	20.672	1.00 39.02
15	ATOM	1375		ARG	840	6.195	15.151	21.731	1.00 40.51
13	ATOM	1376		ARG	840	5.916	16.580	21.828	1.00 42.20
	ATOM	1377		ARG	840	7.032	17.289	22.610	1.00 43.32
	ATOM	1378		ARG	840	6.657	18.639	23.261	1.00 45.47
	ATOM	1379		ARG	840	6.945	19.881	22.401	1.00 46.95
20		1380		ARG	840	8.319	20.371	22.542	1.00 48.57
20	ATOM	1381		ARG	840	9.066	20.823	21.533	1.00 49.57
	ATOM	1382	NH1		840	8.580	20.860	20.294	1.00 49.89
	ATOM	1383	NH2		840	10.314	21.220	21.755	1.00 49.90
	ATOM	1384		ARG	840	5.776	17.220	20.457	1.00 43.27
25	ATOM			ARG	840	4.860	18.004	20.232	1.00 43.50
25	ATOM	1385		ILE	841	6.663	16.876	19.528	1.00 44.53
	ATOM	1386	N		841	6.600	17.483	18.211	1.00 46.22
	ATOM	1387	CA	ILE		7.983	17.572	17.510	1.00 45.82
	ATOM	1388	CB	ILE	841 841	9.044	18.078	18.463	1.00 46.40
00	ATOM	1389		ILE		8.383	16.237	16.918	1.00 46.46
30	ATOM	1390	CG1	ILE	841	8.064	16.150	15.463	1.00 45.92
	MOTA	1391	CD1	ILE	841	5.534	16.913	17.286	1.00 48.04
	MOTA	1392	C	ILE	841	5.472	17.272	16.109	1.00 48.90
	MOTA	1393	0	ILE	841		15.976	17.786	1.00 49.91
	MOTA	1394	N	ILE	842	4.737	15.446	16.990	1.00 51.38
35	MOTA	1395	CA	ILE	842	3.632 3.577	13.446	16.889	1.00 51.11
	ATOM	1396	CB	ILE	842		13.445	15.482	1.00 51.75
	MOTA	1397	CG2	ILE	842	3.917	13.192	17.870	1.00 50.67
	ATOM	1398	CG1	ILE	842	4.523	11.701	17.619	1.00 49.36
4.0	MOTA	1399	CD1	ILE	842	4.691		17.659	1.00 52.63
40	ATOM	1400	C	ILE	842	2.384	16.003 16.551	16.999	1.00 52.38
	MOTA	1401	0	ILE	842	1.509		18.986	1.00 54.81
	MOTA	1402	N	ALA	843	2.356	15.939 16.456	19.761	1.00 57.24
	MOTA	1403	CA	ALA	843	1.242	16.129	21.247	1.00 56.69
	MOTA	1404	CB	ALA	843	1.424		19.557	1.00 59.28
45	MOTA	1405	C	ALA	843	1.215	17.962	20.304	1.00 59.69
	MOTA	1406	0	ALA	843	1.847	18.704		1.00 61.60
	ATOM	1407	N	CYS	844	0.560	18.391	18.481	1.00 63.67
	ATOM	1408	CA	CYS	844	0.402	19.810 20.536	17.979	1.00 64.05
	MOTA	1409	СВ	CYS	844	1.766	20.336	16.470	1.00 65.30
50	MOTA	1410	SG	CYS	844	2.751		16.848	1.00 64.63
	MOTA	1411	C	CYS	844	-0.441	19.854	16.889	1.00 64.70
	MOTA	1412	0	CYS	844	-1.618	19.471	15.738	1.00 65.65
	ATOM	1413	N	ALA	845	0.136	20.332		
	ATOM	1414	CA	ALA	845	-0.545	20.374	14.439	1.00 65.96
55	MOTA	1415	CB	ALA	845	-0.195	21.639	13.684	1.00 65.80
	ATOM	1416	С	ALA	845	-0.079	19.165	13.644	1.00 66.49
	MOTA	1417	0	ALA	845	-0.675	18.829	12.620	1.00 66.85
	MOTA	1418	N	ALA	846	0.998	18.533	14.127	1.00 66.74
	ATOM	1419	CA	ALA	846	1.601	17.343	13.511	1.00 66.90
60	ATOM	1420	CB	ALA	846	3.110	17.337	13.730	1.00 66.74
	MOTA	1421	С	ALA	846	0.984	16.074	14.086	1.00 66.88
	MOTA	1422	0	ALA	846	1.675	15.092	14.345	1.00 66.35

								- 4 201	1.00 67.27
	ATOM	1423	N	ALA	847	-0.325	16.141	14.291	1.00 67.27
	ATOM	1424	CA	ALA	847	-1.162	15.076	14.826	1.00 68.14
	ATOM	1425	СВ	ALA	847	-0.515	14.407	16.033	
	ATOM	1426	С	ALA	847	-2.420	15.816	15.251	
5	ATOM	1427	0	ALA	847	-2.432	17.046	15.278	
J	ATOM	1428	N	ALA	848	-3.468	15.079	15.597	1.00 69.11
	ATOM	1429	CA	ALA	848	-4.728	15.685	16.016	1.00 69.71
	ATOM	1430	CB	ALA	848	-5.272	16.598	14.907	1.00 69.68
		1431	Ċ	ALA	848	-5.737	14.586	16.340	1.00 70.20
10	ATOM	1432	ō	ALA	848	-5.342	13.474	16.720	1.00 70.21
10	ATOM	1433	N	ALA	849	-7.021	14.914	16.146	1.00 70.54
	ATOM			ALA	849	-8.185	14.043	16.374	1.00 70.31
	ATOM	1434	CA	ALA	849	-9.014	13.929	15.079	1.00 70.65
	MOTA	1435	СВ		849	-7.856	12.661	16.920	1.00 69.92
	MOTA	1436	C	ALA	849	-7.665	12.492	18.130	1.00 70.12
15	ATOM	1437	0	ALA		-7.808	11.680	16.020	1.00 69.47
	ATOM	1438	N	ALA	850	-7.473	10.304	16.377	1.00 68.74
	ATOM	1439	CA	ALA	850	-8.494	9.331	15.774	1.00 68.95
	MOTA	1440	CB	ALA	850	-6.061	10.015	15.845	1.00 67.76
	MOTA	1441	С	ALA	850		8.867	15.864	1.00 67.27
20	ATOM	1442	0	ALA	850	-5.590	11.077	15.388	1.00 66.46
	ATOM	1443	N	SER	851	-5.391		14.846	1.00 65.28
	ATOM	1444	CA	SER	851	-4.046	10.962		1.00 65.24
	MOTA	1445	CB	SER	851	-3.664	12.200	14.018	1.00 65.05
	ATOM	1446	OG	SER	851	-2.405	12.037	13.369	1.00 64.16
25	ATOM	1447	С	SER	851	-3.023	10.710	15.944	1.00 64.16
	ATOM	1448	0	SER	851	-2.426	9.639	15.969	1.00 62.68
	ATOM	1449	N	CYS	852	-2.873	11.651	16.879	
	ATOM	1450	CA	CYS	852	-1.901	11.526	17.974	1.00 60.96
	ATOM	1451	CB	CYS	852	-2.353	12.333	19.191	1.00 61.34
30	ATOM	1452	SG	CYS	852	-1.712	14.024	19.216	1.00 62.06
50	ATOM	1453	C	CYS	852	-1.480	10.102	18.388	1.00 59.71
	MOTA	1454	Ö	CYS	852	-0.282	9.800	18.422	1.00 59.67
	MOTA	1455	N	SER	853	-2.440	9.223	18.678	1.00 57.85
	ATOM	1456	CA	SER	853	-2.109	7.847	19.053	1.00 55.45
35	ATOM	1457	СВ	SER	853	-3.308	7.133	19.686	1.00 56.27
33		1458	OG	SER	853	-3.480	7.522	21.038	1.00 57.65
	ATOM	1459	C	SER	853	-1.610	7.052	17.847	1.00 53.09
	MOTA		ō	SER	853	-0.601	6.346	17.942	1.00 53.22
	ATOM	1460	N	ARG	854	-2.304	7.172	16.719	1.00 49.75
40	MOTA	1461		ARG	854	-1.924	6.462	15.500	1.00 46.39
40	ATOM	1462	CA		854	-3.021	6.634	14.443	1.00 47.23
	MOTA	1463	CB	ARG	854	-2.747	5.970	13.105	1.00 48.09
	MOTA	1464	CG	ARG	854	-2.985	6.943	11.940	1.00 49.85
	ATOM	1465	CD	ARG		-4.340	7.500	11.936	1.00 51.11
	ATOM	1466	NE	ARG	854	-4.911	8.122	10.904	1.00 51.20
45	MOTA	1467	CZ	ARG	854	-4.261	8.290	9.755	1.00 51.32
	MOTA	1468		LARG	854		8.567	11.021	1.00 51.21
	MOTA	1469		2 ARG	854	-6.153	6.962	14.967	1.00 43.72
	MOTA	1470	С	ARG	854	-0.564	6.164	14.564	1.00 43.80
	ATOM	1471.		ARG	854	0.294		15.019	1.00 39.68
50	ATOM	1472	N	ARG	855	-0.360		14.558	1.00 36.58
	ATOM	1473	CA	ARG	855	0.860			1.00 36.42
	ATOM	1474	CB	ARG	855	0.713		14.734	1.00 34.79
	ATOM	1475	CG	ARG	855	1.801		14.082	
	MOTA	1476	CD	ARG	855	1.724			1.00 32.86
55	ATOM	1477	NE	ARG	855	2.869			1.00 30.54
	ATOM	1478	CZ	ARG	855	3.030			1.00 29.61
	ATOM	1479		1 ARG	855	2.130		9.831	1.00 28.88
	ATOM	1480		2 ARG	855	4.085			1.00 29.20
	ATOM	1481		ARG	855	2.106	8.454		1.00 35.02
60	ATOM	1482		ARG	855	3.180	8.326	14.723	1.00 34.97
JU	MOTA	1483		PHE	856	1.973		16.609	
	ATOM	1484			856	3.086			1.00 31.21
	AION	7101							

	ATOM	1485	CB	PHE	856	2.846	8.128	18.895	1.00 30.24
	ATOM	1486	CG	PHE	856	4.058	7.977	19.755	1.00 30.06
	ATOM	1487	CD1	PHE	856	5.022	8.972	19.787	1.00 29.34
	ATOM	1488	CD2	PHE	856	4.241	6.829	20.536	1.00 29.33
5	ATOM	1489		PHE	856	6.150	8.832	20.580	1.00 29.36
9	ATOM	1490		PHE	856	5.364	6.676	21.335	1.00 28.86
	ATOM	1491		PHE	856	6.325	7.680	21.357	1.00 29.62
		1492		PHE	856	3.308	6.290	17.254	1.00 30.61
	ATOM	1493		PHE	856	4.420	5.803	17.424	1.00 30.75
40	ATOM	1494		TYR	857	2.258	5.543	16.943	1.00 29.79
10	ATOM		-	TYR	857	2.446	4.118	16.725	1.00 29.79
	ATOM	1495		TYR	857	1.116	3.365	16.660	1.00 30.78
	MOTA	1496			857	1.254	1.871	16.396	1.00 32.21
	MOTA	1497		TYR		1.425	0.972	17.442	1.00 32.75
	MOTA	1498		TYR	857	1.548	-0.401	17.215	1.00 34.60
15	MOTA	1499		TYR	857		1.362	15.098	1.00 33.36
	ATOM	1500		TYR	857	1.208	-0.011	14.854	1.00 34.67
	ATOM	1501		TYR	857	1.331		15.918	1.00 35.61
	ATOM	1502		TYR	857	1.503	-0.887	15.697	1.00 36.96
	MOTA	1503		TYR	857	1.652	-2.244		1.00 29.44
20	MOTA	1504		TYR	857	3.206	3.929	15.419	1.00 29.44
	MOTA	1505	0	TYR	857	4.135	3.125	15.371	1.00 29.44
	MOTA	1506	N	GLN	858	2.847	4.685	14.376	
	ATOM	1507	CA	GLN	858	3.533	4.537	13.087	1.00 28.06
	ATOM	1508	CB	GLN	858	2.675	5.020	11.890	1.00 28.98
25	ATOM	1509	CG	GLN	858	1.970	6.384	12.029	1.00 31.65
	ATOM	1510	CD	GLN	858	0.781	6.569	11.059	1.00 32.26
	ATOM	1511	OE1	GLN	858	0.385	7.700	10.724	1.00 32.11
	ATOM	1512	NE2		858	0.210	5.458	10.617	1.00 32.79
	ATOM	1513	С	GLN	858	4.967	5.077	13.037	1.00 26.56
30	ATOM	1514	Ö	GLN	858	5.820	4.525	12.332	1.00 26.58
50	ATOM	1515	N	LEU	859	5.266	6.101	13.825	1.00 24.64
	ATOM	1516	CA	LEU	859	6.622	6.632	13.832	1.00 22.82
	ATOM	1517	CB	LEU	859	6.675	8.067	14.395	1.00 23.93
	ATOM	1518	CG	LEU	859	6.054	9.243	13.617	1.00 23.18
35	ATOM	1519	CD1		859	6.616	10.541	14.156	1.00 23.42
55	ATOM	1520	CD2		859	6.363	9.137	12.173	1.00 22.30
	ATOM	1521	C	LEU	859	7.545	5.705	14.613	1.00 20.92
		1522	Ö	LEU	859	8.694	5.486	14.222	1.00 20.35
	ATOM	1523	N	THR	860	7.030	5.124	15.691	1.00 20.30
40	ATOM	1524	CA	THR	860	7.821	4.195	16.505	1.00 20.14
40	ATOM		CB	THR	860	7.215	3.957	17.905	1.00 18.65
	ATOM	1525		THR	860	5.849	3.551	17.797	1.00 18.55
	MOTA	1526	OG1	THR	860	7.314	5.196	18.734	1.00 18.12
	ATOM	1527	CG2	THR	860	7.969	2.855	15.765	1.00 20.30
45	ATOM	1528	C	THR	860	8.922	2.108	15.985	1.00 20.28
45	ATOM	1529	0		861	7.040	2.600	14.851	1.00 21.02
	MOTA	1530	N	LYS	861	7.046	1.411	14.034	1.00 21.82
	ATOM	1531	CA	LYS		5.649	1.178	13.475	1.00 23.82
	ATOM	1532	CB	LYS	861	5.375	-0.268	13.110	1.00 26.71
	ATOM	1533	CG	LYS	861		-1.106	14.321	1.00 28.40
50	MOTA	1534	CD	LYS	861	5.015		13.922	1.00 20.40
	MOTA	1535	CE	LYS	861	4.924	-2.587		1.00 30.14
	MOTA	1536	NZ	LYS	861	4.542	-3.514	15.050	1.00 31.33
	ATOM	1537	С	LYS	861	8.040	1.642	12.895	
	MOTA	1538	0	LYS	861	8.781	0.750	12.510	1.00 21.01
55	MOTA	1539	N	LEU	862	8.093	2.863	12.389	1.00 21.49
	ATOM	1540	CA	LEU	862	9.021	3.192	11.311	1.00 21.32
	MOTA	1541	CB	LEU	862	8.713	4.577	10.731	1.00 22.14
	MOTA	1542	CG	LEU	862	9.816	5.281	9.921	1.00 21.98
	ATOM	1543		LEU	862	10.022	4.563	8.622	1.00 22.72
60	ATOM	1544	CD2	LEU	862	9.456	6.726	9.670	1.00 22.58
	ATOM	1545	С	LEU	862	10.438	3.181	11.854	1.00 21.15
	ATOM	1546	0	LEU	862	11.386	2.884	11.133	1.00 22.30

	ATOM ATOM	1547 1548	-	LEU LEU	863 863	1	10.596	3.602 3.625	13.105 13.749 15.040	1.00 20.71 1.00 18.90 1.00 18.79
	ATOM	1549		LEU	863		1.827	4.423 5.931	14.863	1.00 18.02
	MOTA	1550		LEU	863		11.890 12.103	6.545	16.230	1.00 19.67
5	ATOM	1551	CD1		863		13.049	6.291	13.944	1.00 16.59
	MOTA	1552	CD2		863		12.462	2.218	14.004	1.00 18.08
	MOTA	1553	-	LEU	863 863		13.676	1.984	13.895	1.00 17.06
	ATOM	1554	-	LEU ASP	864		11.592	1.307	14.436	1.00 18.33
40	ATOM	1555		ASP	864		11.985	-0.088	14.642	1.00 19.28
10	ATOM	1556		ASP	864		10.797	-0.917	15.143	1.00 19.27
	ATOM	1557 1558		ASP	864		10.525	-0.727	16.620	1.00 19.92
	ATOM	1559	OD1		864		11.256	0.045	17.271	1.00 20.94
	ATOM ATOM	1560	OD2		864		9.577	-1.364	17.116	1.00 19.39
15	ATOM	1561		ASP	864		12.467	-0.692	13.321	1.00 19.06
13	ATOM	1562	_	ASP	864		13.377	-1.519	13.298	1.00 18.82
	ATOM	1563		SER	865		11.847	-0.263	12.222	1.00 19.90
	ATOM	1564	CA	SER	865		12.202	-0.764	10.894	1.00 19.17
	ATOM	1565	CB	SER	865		11.226	-0.289	9.798	1.00 18.59 1.00 19.91
20	ATOM	1566		SER	865		11.167	1.123	9.613	1.00 19.91 1.00 18.01
	ATOM	1567	С	SER	865		13.634	-0.507	10.489 9.765	1.00 18.93
	MOTA	1568	0	SER	865		14.213	-1.294 0.535	11.004	1.00 17.08
	MOTA	1569	N	VAL	866		14.257 15.619	0.747	10.589	1.00 15.20
	MOTA	1570	CA	VAL	866		16.093	2.211	10.783	1.00 15.01
25	MOTA	1571	CB	VAL	866		14.982	3.081	11.320	1.00 13.81
	ATOM	1572		VAL	866 866		17.344	2.280	11.574	1.00 13.41
	ATOM	1573	CG2	VAL	866		16.564	-0.260	11.194	1.00 14.83
	ATOM	1574	С 0	VAL	866		17.625	-0.518	10.641	1.00 14.66
30	ATOM	1575 1576	N	GLN	867		16.168	-0.873	12.302	1.00 15.12
30	ATOM ATOM	1577	CA	GLN	867		17.031	-1.849	12.977	1.00 15.64
	ATOM	1578	CB	GLN	867		16.508	-2.155	14.374	1.00 16.10
	ATOM	1579	ĊĠ	GLN	867		16.526	-0.968	15.315	1.00 16.35
	ATOM	1580	CD	GLN	867		17.910	-0.474	15.672	1.00 17.91
35	ATOM	1581		GLN	867		18.924	-1.175	15.510	1.00 17.76
	MOTA	1582	NE2	GLN	867		17.958	0.750	16.201	1.00 17.53 1.00 14.54
	MOTA	1583	С	GLN	867		17.358	-3.143	12.233	1.00 14.54 1.00 15.92
	MOTA	1584	0	GLN	867		18.487	-3.594	12.271 11.634	1.00 13.32
	MOTA	1585	N	PRO	868		16.364	-3.809 -3.555	11.696	1.00 15.17
40	ATOM	1586	CD	PRO	868		14.914	-5.040	10.886	1.00 13.73
	MOTA	1587	CA	PRO	868		16.630 15.232	-5.465	10.415	1.00 14.35
	MOTA	1588	CB	PRO	868 868		14.331	-4.928	11.438	1.00 14.42
	ATOM	1589	CG	PRO PRO	868		17.500	-4.704	9.674	1.00 13.45
45	ATOM	1590 1591	C O	PRO	868		18.341	-5.497	9.254	1.00 14.77
43	ATOM ATOM	1592	N	ILE	869		17.289	-3.514	9.113	1.00 13.49
	ATOM	1593	CA	ILE	869		18.043	-3.044	7.970	1.00 12.06
	ATOM	1594	СВ	ILE	869		17.447	-1.740	7.358	1.00 12.53
	ATOM	1595	CG2		869		18.272		6.175	1.00 12.44
50	ATOM	1596	CG1		869		15.998		6.928	1.00 12.25
••	ATOM	1597	CD1	ILE	869		15.258		6.432	1.00 11.91
	ATOM	1598	С	ILE	869		19.458		8.411	1.00 11.70
	MOTA	1599	0	ILE	869		20.356		7.755	1.00 13.01
	ATOM	1600	N	ALA	870		19.655		9.610	1.00 12.19 1.00 11.52
55	ATOM	1601	CA	ALA	870		21.007		10.110	1.00 11.52
	MOTA	1602	CB	ALA	870		20.971			1.00 10.82
	MOTA	1603	C	ALA	870		21.758		10.350	1.00 12.90
	ATOM	1604	0	ALA	870		22.955			1.00 13.99
00	ATOM	1605	N	ARG	871		21.082 21.659			1.00 15.30
60	ATOM	1606	CA	ARG	871 871		20.668			1.00 16.99
	ATOM	1607	CB	ARG ARG	871 871		21.317			1.00 20.44
	MOTA	1608	ÇG	ANG	0 / 1			. , , , , ,		

	ATOM ATOM	1609 1610		ARG ARG	871 871	20.552 21.529	-8.755 -9.736	13.190 13.678	1.00 22.19 1.00 25.05
	ATOM	1611		ARG	871	22.248	-9.581	14.785	1.00 24.87
	ATOM	1612	NH1		871	22.085	-8.513	15.553	1.00 26.86
5	ATOM	1613	NH2		871	23.221		15.059	1.00 27.12
3	ATOM	.1614		ARG	871	22.119	-6.287	9.939	1.00 16.18
	ATOM	1615		ARG	871	23.216	-6.846	9.897	1.00 16.90
	ATOM	1616		GLU	872	21.300	-6.256	8.886	1.00 17.08
	ATOM	1617		GLU	872	21.669	-6.874	7.595	1.00 17.70
10	ATOM	1618		GLU	872	20.546	-6.670	6.578	1.00 20.21
10	ATOM	1619		GLU	872	20.070	-7.920	5.827	1.00 27.32
	ATOM	1620		GLU	872	19.041	-7.600	4.715	1.00 31.24
	ATOM	1621	OE1	GLU	872	19.199	-8.069	3.544	1.00 32.65
	ATOM	1622	OE2		872	18.068	-6.867	5.018	1.00 33.14
15	ATOM	1623	С	GLU ·	872	22.961	-6.229	7.064	1.00 16.22
	ATOM	1624		GLU	872	23.826	-6.892	6.504	1.00 16.64
	ATOM	1625	N	LEU	873	23.109	-4.927	7.254	1.00 15.48
	ATOM	1626	CA	LEU	873	24.304	-4.230	6.781	1.00 13.64
	ATOM	1627	CB	LEU	873	24.040	-2.718	6.664	1.00 13.09
20	ATOM	1628	CG	LEU	873	22.957	-2.359	5.640	1.00 12.60
	ATOM	1629	CD1	LEU	873	22.396	-0.985	5.856	1.00 13.12
	ATOM	1630	CD2	LEU	873	23.511	-2.529	4.229	1.00 12.29
	ATOM	1631	С	LEU	873	25.489	-4.510	7.662	1.00 13.26
	ATOM	1632	0	LEU	873	26.621	-4.541	7.185	1.00 12.91 1.00 14.96
25	ATOM	1633	N	HIS	874	25.237	-4.688	8.960	1.00 14.98
	ATOM	1634	CA	HIS	874	26.297	-5.011	9.935	1.00 13.73
	MOTA	1635	CB	HIS	874	25.735	-5.154	11.351	1.00 14.03
	ATOM	1636	CG	HIS	874	25.513	-3.860	12.062 12.204	1.00 13.33
	MOTA	1637	CD2		874	26.303	-2.769	12.204	1.00 12.74
30	MOTA	1638		HIS	874	24.365	-3.588	13.313	1.00 12.74
	MOTA	1639	CE1		874	24.451	-2.397	12.990	1.00 10.87
	ATOM	1640	NE2		874	25.616	-1.878 -6.342	9.549	1.00 16.66
	MOTA	1641	С	HIS	874	26.945	-6.454	9.539	1.00 16.68
	ATOM	1642	0	HIS	874	28.171	-7.356	9.268	1.00 18.67
35	MOTA	1643	N	GLN	875	26.122	-8.674	8.853	1.00 19.45
	ATOM	1644	CA	GLN	875	26.635 25.507	-9.726	8.779	1.00 21.56
	ATOM	1645	СВ	GLN	875		-10.875	9.864	1.00 25.76
	MOTA	1646	CG	GLN	875		-11.938	9.671	1.00 26.74
4.0	ATOM	1647	CD	GLN	875		-11.624	9.654	1.00 27.36
40	ATOM	1648	OE1	GLN	875	26.285	-13.204	9.589	1.00 27.93
	ATOM	1649		GLN	875 875	27.324	-8.521	7.491	1.00 18.37
	ATOM	1650	С	GLN GLN	875	28.428	-9.022	7.294	1.00 18.65
	ATOM	1651	0	PHE	876	26.737	-7.724	6.597	1.00 18.47
45	ATOM	1652 1653	N CA	PHE	876	27.338	-7.515	5.280	1.00 18.22
40	ATOM ATOM	1654	CB	PHE	876	26.453	-6.641	4.377	1.00 19.25
		1655	CG	PHE	876	26.966	-6.506	2.954	1.00 19.63
	ATOM ATOM	1656		PHE	876	28.038	-5.675	2.657	1.00 18.97
	ATOM	1657		PHE	876	26.380	-7.226	1.917	1.00 19.90
50	ATOM	1658		PHE	876	28.519	-5.558	1.343	1.00 20.30
50	ATOM	1659		PHE	876	26.857	-7.113	0.597	1.00 20.70
	ATOM	1660	CZ	PHE	876	27.926	-6.281	0.310	1.00 18.82
	ATOM	1661	c	PHE	876	28.689	-6.871	5.403	1.00 17.76
	ATOM	1662	ŏ	PHE	876	29.687	-7.412	4.920	1.00 17.95
55	ATOM	1663	N	THR	877	28.741	-5.732	6.086	1.00 17.85
00	MOTA	1664	CA	THR	877	30.002	-5.024	6.215	1.00 17.77
	ATOM	1665	CB	THR	877	29.855		6.915	1.00 18.24
	ATOM	1666		THR	877	30.954		6.525	1.00 19.13
	ATOM	1667	CG2		877	29.868		8.444	1.00 17.92
60	ATOM	1668	С	THR	877	31.040		6.900	1.00 17.52
	ATOM	1669	0	THR	877	32.208		6.514	1.00 16.51
	ATOM	1670	N	PHE	878	30.634	-6.610	7.943	1.00 18.06

	ATOM ATOM	1671 1672		HE	878 878	31.559 30.863	-7.501 -8.201	9.805	1.00 19.20 1.00 19.53
	ATOM	1673		PHE	878	31.731	-9.220	10.484	1.00 20.60
	ATOM	1674	CD1 E		878	32.681	-8.829	11.414	1.00.19.88
5	ATOM	1675	CD2 F		878	31.623		10.150	1.00 20.59
9	ATOM	1676	CE1 E		878	33.518	-9.774	12.008	1.00 22.10
	ATOM	1677	CE2		878	32.454	-11.532	10.733	1.00 20.21
	ATOM	1678		PHE	878	33.403	-11.138	11.660	1.00 20.82
	ATOM	1679		PHE	878	32.176	-8.567	7.725	1.00 18.91
10		1680	-	PHE	878	33.400	-8.724	7.670	1.00 17.63
10	ATOM	1681		ASP	879	31.326	-9.268	6.973	1.00 19.57
	ATOM	1682	-	ASP	879	31.800	-10.301	6.054	1.00 20.02
	ATOM	1683		ASP	879	30.622	-10.972	5.342	1.00 20.24
	ATOM	1684		ASP	879	29.693	-11.724	6.307	1.00 22.04
4 =	ATOM	1685	OD1		879	30.122	-12.072	7.443	1.00 23.16
15	MOTA		OD2		879	28.520	-11.968	5.937	1.00 21.98
	MOTA	1686		ASP	879	32.723	-9.654	5.044	1.00 20.35
	ATOM	1687		ASP	879	33.802	-10.171	4.737	1.00 20.51
	ATOM	1688		LEU	880	32.342	-8.472	4.580	1.00 20.77
00	MOTA	1689		LEU	880	33.149	-7.775	3.596	1.00 20.33
20	ATOM	1690	-	LEU	880	32.484	-6.471	3.180	1.00 20.23
	ATOM	1691		LEU	880	33.089	-5.838	1.939	1.00 18.50
	MOTA	1692			880	33.310	-6.886	0.855	1.00 19.38
	MOTA	1693	CD1		880	32.159	-4.762	1.477	1.00 18.07
	MOTA	1694	CD2		880	34.529	-7.496	4.136	1.00 20.40
25	MOTA	1695	-	LEU		35.513	-7.723	3.453	1.00 21.41
	ATOM	1696		LEU	880	34.602	-7.040	5.376	1.00 20.90
	MOTA	1697	-	LEU	881 881	35.882	-6.723	6.011	1.00 20.84
	ATOM	1698		LEU		35.651	-6.055	7.364	1.00 19.23
	MOTA	1699		LEU	881	36.989	-5.773	8.031	1.00 19.26
30	MOTA	1700		LEU	881	37.662	-4.593	7.350	1.00 19.67
	MOTA	1701	CD1		881	36.810	-5.514	9.500	1.00 18.92
	MOTA	1702	CD2		881	36.818	-7.923	6.188	1.00 21.55
	ATOM	1703		LEU	881	38.055	-7.806	6.107	1.00 21.03
	MOTA	1704		LEU	881	36.230		6.492	1.00 22.39
35	ATOM	1705	N	ILE	882		-10.265	6.671	1.00 23.63
	MOTA	1706	CA	ILE	882	37.013	-11.390	7.248	1.00 23.00
	MOTA	1707	CB	ILE	882	30.130	-12.729	7.185	1.00 22.75
	ATOM	1708	CG2	ILE	882	36.633	-11.006	8.675	1.00 22.91
	MOTA	1709		ILE	882	35.749	-10.412	9.491	1.00 22.94
40	ATOM	1710	CD1	ILE	882		-10.412	5.340	1.00 24.48
	ATOM	1711	С	ILE	882	37.000	-10.043	5.290	1.00 24.23
	ATOM	1712	0	ILE	882	38.839	-10.533	4.256	1.00 25.73
	ATOM	1713	N	LYS	883			2.945	1.00 28.25
	ATOM	1714	CA	LYS	883	37.441	-10.868	2.211	1.00 27.47
45	MOTA	1715	СВ	LYS	883		-11.820	1.932	1.00 27.17
	ATOM	1716	CG	LYS	883	35.140	-11.240	1.109	
	MOTA	1717	CD	LYS	883	34.293	-12.163		1.00 27.00
	ATOM	1718	CE	LYS	883	32.926	-11.544	0.899	1.00 29.99
	ATOM	1719	NZ	LYS	883		-12.319	-0.003 2.061	1.00 29.99
50	ATOM	1720	С	LYS	883	37.749			
	MOTA	1721	0	LYS	883	37.823		0.841	1.00 30.81
	ATOM	1722	N	SER	884	37.976		2.672	1.00 32.24
	ATOM	1723	CA	SER	884	38.268		1.938	1.00 33.66
	ATOM	1724	CB	SER	884	38.440		2.921	1.00 32.96
55	ATOM	1725	0G	SER	884	39.466		3.856	1.00 32.02
	ATOM	1726	С	SER	884	39.500		1.042	1.00 35.48
	ATOM	1727		SER	884	39.491		-0.087	1.00 35.08
	MOTA	1728		HIS	885	40.55		1.556	1.00 38.15
	ATOM	1729		HIS	885	41.815		0.824	1.00 40.84
60	ATOM	1730		HIS	885	42.882		1.789	1.00 43.70
J -	ATOM	1731		HIS	885	44.032	2 -9.392	1.124	1.00 47.44
	ATOM	1732		HIS	885	44.240	-10.707	0.860	1.00 49.14

							0 707	0.704	1.00 49.20
	ATOM	1733	ND1		885	45.172	-8.737	0.704 0.217	1.00 49.74
	ATOM	1734	CE1		885	46.034	-9.615	0.300	1.00 50.17
	ATOM	1735	NE2	HIS	885		10.818		1.00 30.17
	MOTA	1736	C	HIS	885	41.682	-9.017	-0.432	1.00 41.51
5	ATOM	1737	0	HIS	885	42.563	-9.010	-1.288	
•	ATOM	1738	N I	MET	886	40.586	-9.762	-0.544	
	ATOM	1739	CA	MET	886	40.372 -	-10.639	-1.686	1.00 41.17 1.00 43.08
	ATOM	1740	СВ	MET	886	39.859 -	-11.989	-1.212	1.00 45.00
	ATOM	1741	CG	MET	886	40.928	-12.860	-0.584	1.00 45.59
10	ATOM	1742		MET	886	40.175	-14.113	0.457	1.00 50.78
10	ATOM	1743		MET	886	39.069 ·	-14.971	-0.725	1.00 48.51
	ATOM	1744		MET	886	39.455	-10.074	-2.761	1.00 40.68
	ATOM	1745		MET	886	39.535	-10.476	-3.923	1.00 41.66
	ATOM	1746	-	VAL	887	38.542	-9.193	-2.370	1.00 39.25
15	ATOM	1747		VAL	887	37.637	-8.565	-3.333	1.00 37.55
15		1748		VAL	887	36.187	-8.459	-2.802	1.00 37.19
	ATOM	1749	-	VAL	887	35.526	-9.828	-2.756	1.00 37.49
	MOTA	1750		VAL	887	36.175	-7.817	-1.429	1.00 36.99
	ATOM		C	VAL	887	38.145	-7.168	-3.702	1.00 37.08
00	ATOM	1751	0	VAL	887	37.484	-6.444	-4.442	1.00 37.26
20	ATOM	1752		SER	888	39.320	-6.809	-3.188	1.00 35.90
	ATOM	1753	N	SER	888	39.955	-5.515	-3.437	1.00 35.05
	ATOM	1754	CA	SER	888	40.231	-5.342	-4.929	1.00 35.29
	MOTA	1755	CB		888	41.335	-6.133	-5.326	1.00 36.74
~=	MOTA	1756	OG	SER		39.216	-4.290	-2.898	1.00 34.27
25	MOTA	1757	С	SER	888	39.402	-3.179	-3.396	1.00 34.78
	MOTA	1758	0	SER	888	38.391	-4.485	-1.875	1.00 32.78
	MOTA	1759	N	VAL	889	37.636	-3.386	-1.283	1.00 31.50
	MOTA	1760	CA	VAL	889	36.244	-3.857	-0.772	1.00 30.79
	MOTA	1761	СВ	VAL	889	35.509	-2.729	-0.055	1.00 30.12
30	MOTA	1762	CG1		889	_	-4.364	-1.903	1.00 30.08
	ATOM	1763		VAL	889	35.410	-3.002	-0.064	1.00 31.36
	ATOM	1764	С	VAL	889	38.410	-3.895	0.648	1.00 32.20
	ATOM	1765	0	VAL	889	38.855	-1.724	0.156	1.00 31.10
	ATOM	1766	N	ASP	890	38.692	-1.428	1.414	1.00 30.80
35	MOTA	1767	CA	ASP	890	39.364	-1.093	1.296	1.00 33.89
	MOTA	1768	CB	ASP	890	40.849	-1.949	2.261	1.00 35.96
	MOTA	1769	CG	ASP	890	41.720	-2.314	3.373	1.00 35.86
	ATOM	1770		ASP	890	41.248	-2.260	1.901	1.00 37.33
	ATOM	1771		ASP	890	42.882		2.326	1.00 28.51
40	MOTA	1772	С	ASP	890	38.629	-0.493 0.379	1.889	1.00 27.96
	ATOM	1773	0	ASP	890	37.889		3.610	1.00 26.20
	ATOM	1774	N	PHE	891	38.761	-0.782	4.661	1.00 24.15
	ATOM	1775	ÇA	PHE	891	38.096	-0.045	5.732	1.00 20.51
	ATOM	1776	CB	PHE	891	37.595	-1.027	5.259	1.00 16.33
45	ATOM	1777	CG	PHE	891	36.501	-1.937	4.288	1.00 15.42
	ATOM	1778	CD1	PHE	891	36.741	-2.892		1.00 14.37
	ATOM	1779		PHE	891	35.230	-1.826	5.773	1.00 13.60
	MOTA	1780	CE1		891	35.720	-3.730	3.832	1.00 14.04
	ATOM	1781		PHE	891	34.220	-2.648	5.335	1.00 13.43
50	ATOM	1782	CZ	PHE	891	34.467	-3.607	4.353	1.00 24.58
	ATOM	1783	С	PHE	891	39.036	0.942	5.305	1.00 25.30
	ATOM	1784	0	PHE	891	40.150	0.574	5.695	1.00 23.30
	MOTA	1785	N	PRO	892	38.603	2.209	5.437	
	ATOM	1786	CD	PRO	892	37.376	2.811	4.909	1.00 22.44
55	ATOM	1787	ÇA	PRO	892	39.441	3.234	6.060	1.00 23.41
	ATOM	1788	CB	PRO	892	38.582	4.485	5.940	1.00 23.21
	ATOM	1789	CG	PRO	892	37.796	4.241	4.748	1.00 23.19
	ATOM	1790	С	PRO	892	39.655	2.866	7.520	1.00 23.31
	ATOM	1791	0	PRO	892	38.887	2.090	8.078	1.00 22.72
60	ATOM	1792	N	GLU	893	40.619	3.517	8.157	1.00 24.55
J	ATOM	1793	CA	GLU	893	40.984	3.267	9.555	1.00 26.50
	ATOM	1794	CB	GLU	893	41.885	4.385	10.072	1.00 28.90
	•								

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								11 500	1.00 33.98
	ATOM	1795	CG	GLU	893	42.329	4.192	11.509	1.00 33.38
	ATOM	1796	CD	GLU	893	42.441	5.498	12.280	1.00 37.50
	ATOM	1797	OE1	GLU	893	43.356	6.292	11.955	1.00 39.34
	ATOM	1798	OE2	GLU	893	41.624	5.729	13.216	1.00 39.34
5	ATOM	1799	С	GLU	893	39.859	3.054	10.563	
•	ATOM	1800	0	GLU	893	39.750	1.992	11.180	1.00 27.29
	ATOM	1801	N	MET	894	39.052	4.078	10.782	1.00 26.07
	ATOM	1802	CA	MET	894	37.968	3.974	11.744	1.00 26.28
	ATOM	1803	CB	MET	894	37.313	5.337	11.954	1.00 28.30
10	ATOM	1804	ĊĠ	MET	894	38.256	6.389	12.509	1.00 32.56
10	MOTA	1805	SD	MET	894	38.847	5.925	14.144	1.00 38.01
	ATOM	1806	CE	MET	894	37.260	5.830	15.037	1.00 35.95
	ATOM	1807	c	MET	894	36.927	2.918	11.393	1.00 24.69
		1808	õ	MET	894	36.337	2.311	12.287	1.00 24.64
46	ATOM	1809	N	MET	895	36.662	2.743	10.102	1.00 23.64
15	ATOM		CA	MET	895	35.705	1.738	9.645	1.00 22.83
	MOTA	1810		MET	895	35.487	1.824	8.135	1.00 21.32
	ATOM	1811	CB		895	34.669	3.006	7.693	1.00 21.17
	ATOM	1812	CG	MET	895	33.044	3.064	8.432	1.00 20.56
	MOTA	1813	SD	MET		32.088	2.305	7.205	1.00 22.81
20	MOTA	1814	CE	MET	895	36.171	0.328	10.032	1.00 22.26
	MOTA	1815	С	MET	895	35.469	-0.383	10.714	1.00 22.26
	MOTA	1816	0	MET	895		-0.066	9.616	1.00 22.06
	MOTA	1817	N	ALA	896	37.362	-1.378	9.953	1.00 22.36
	MOTA	1818	CA	ALA	896	37.867	-1.588	9.350	1.00 22.56
25	MOTA	1819	CB	ALA	896	39.243	-1.581	11.460	1.00 22.96
	MOTA	1820	С	ALA	896	37.914		11.947	1.00 23.87
	MOTA	1821	0	ALA	896	37.520	-2.630	12.212	1.00 23.92
	ATOM	1822	N	GLU	897	38.377	-0.586		1.00 23.32
	ATOM	1823	CA	GLU	897	38.455	-0.724	13.666	1.00 24.03
30	ATOM	1824	CB	GLU	897	39.128	0.502	14.313	
	ATOM	1825	CG	GLU	897	39.288	0.390	15.841	
	ATOM	1826	CD	GLU	897	39.150	1.718	16.555	1.00 27.88
	MOTA	1827	OE1	GLU	897	40.150	2.453	16.674	1.00 29.49
	ATOM	1828	OE2	GLU	897	38.036	2.018	17.013	1.00 29.22
35	ATOM	1829	С	GLU	897	37.076	-0.901	14.276	1.00 22.80
••	ATOM	1830	0	GLU	897	36.873	-1.774	15.094	1.00 22.95
	ATOM	1831	N	ILE	898	36.129	-0.071	13.884	1.00 22.19
	ATOM	1832	CA	ILE	898	34.801	-0.178	14.459	1.00 21.88
	ATOM	1833	СВ	ILE	898	33.940	1.077	14.196	1.00 21.85
40	ATOM	1834	CG2	ILE	898	32.478	0.836	14.587	1.00 22.66
	ATOM	1835	CG1		898	34.438	2.233	15.043	1.00 22.82
	ATOM	1836	CDI		898	33.490	3.390	15.019	1.00 23.11
	ATOM	1837	C	ILE	898	34.080	-1.398	13.968	1.00 20.49
	ATOM	1838	ō	ILE	898	33.228	-1.917	14.656	1.00 21.90
45	ATOM	1839	N	ILE	899	34.410	-1.860	12.781	1.00 19.59
40	ATOM	1840	CA	ILE	899	33.747	-3.027	12.248	1.00 19.42
	ATOM	1841	CB	ILE	899	33.758	-3.014	10.706	1.00 19.12
	ATOM	1842		2 ILE	899	33.095	-4.285	10.157	1.00 18.71
	ATOM	1843		ILE	899	32.987	-1.786	10.187	1.00 18.56
50	ATOM	1844		l ILE	899	33.054	-1.588	8.683	1.00 15.05
50		1845	C C	ILE	899	34.305	-4.338	12.832	1.00 19.02
	ATOM	1846	Ö	ILE	899	33.571	-5.300	12.982	1.00 19.98
	ATOM	1847	Ŋ	SER	900	35.565	-4.344	13.233	1.00 19.03
	ATOM			SER	900	36.177	-5.518	13.822	1.00 19.74
55	ATOM	1848	CA CB	SER	900	37.614	-5.631	13.340	1.00 19.62
55	ATOM	1849		SER	900	38.368	-4.478	13.683	1.00 22.08
	ATOM	1850	OG		900	36.135	-5.502	15.355	1.00 20.48
	ATOM	1851	С	SER	900	36.352	-6.521	16.010	1.00 21.19
	ATOM	1852	0	SER		35.866	-4.346	15.939	1.00 20.99
60	ATOM	1853		VAL	901 901	35.808	-4.235	17.396	
60	ATOM	1854	CA		901	36.705	-3.074	17.927	1.00 20.22
	ATOM	1855			901	36.407	-2.785	19.382	
	MOTA	1856	CG	1 VAL	301	50.407	203	13.502	

							2 426	17 702	1.00 18.81
	ATOM	1857	CG2 '		901	38.168	-3.436	17.782	1.00 18.31
	ATOM	1858	C '	VAL	901	34.397	-4.087	17.935	1.00 20.42
	MOTA	1859	0 '	VAL	901	33.999	-4.823	18.841	
	MOTA	1860	N	GLN	902	33.614	-3.187	17.350	1.00 19.34
5	ATOM	1861	CA	GLN	902	32.264	-2.957	17.828	1.00 17.55
•	ATOM	1862	CB	GLN	902	31.929	-1.476	17.735	1.00 19.32
	ATOM	1863	CG	GLN	902	32.952	-0.579	18.371	1.00 20.82
	ATOM	1864		GLN	902	33.089	-0.776	19.861	1.00 23.15
	ATOM	1865		GLN	902	32.211	-1.336	20.528	1.00 23.22
10	ATOM	1866		GLN	902	34.197	-0.288	20.404	1.00 25.36
.0	ATOM	1867		GLN	902	31.145	-3.766	17.207	1.00 16.24
	ATOM	1868	-	GLN	902	30.337	-4.326	17.938	1.00 15.40
		1869		VAL	903	31.075	-3.810	15.872	1.00 15.79
	MOTA			VAL	903	30.025	-4.552	15.144	1.00 15.22
4 6	MOTA	1870		VAL	903	30.195	-4.461	13.594	1.00 14.30
15	ATOM	1871	-		903	29.159	-5.314	12.883	1.00 13.20
	MOTA	1872		VAL		30.012	-3.005	13.147	1.00 14.90
	ATOM	1873		VAL	903		-6.010	15.605	1.00 14.74
	ATOM	1874		VAL	903	29.860	-6.489	15.693	1.00 14.48
	MOTA	1875		VAL	903	28.732		15.893	1.00 14.65
20	ATOM	1876		PRO	904	30.976	-6.729	15.571	1.00 13.72
	MOTA	1877		PRO	904	32.377	-6.425		1.00 15.72
	ATOM	1878		PRO	904	30.884	-8.122	16.356	
	ATOM	1879	CB	PRO	904	32.350	-8.481	16.602	1.00 15.45
	ATOM	1880	CG	PRO	904	33.014	-7.830	15.512	1.00 14.60
25	MOTA	1881	С	PRO	904	30.053	-8.206	17.632	1.00 16.77
	ATOM	1882	0	PRO	904	29.151	-9.039	17.713	1.00 18.38
	ATOM	1883	N	LYS	905	30.286	-7.295	18.589	1.00 17.00
	ATOM	1884	CA	LYS	905	29.525	-7.292	19.830	1.00 16.34
	ATOM	1885		LYS	905	29.866	-6.085	20.668	1.00 18.17
30	ATOM	1886		LYS	905	31.293	-6.007	21.132	1.00 19.96
00	ATOM	1887		LYS	905	31.464	-4.733	21.947	1.00 22.09
	ATOM	1888		LYS	905	32.911	-4.429	22.276	1.00 23.59
	ATOM	1889		LYS	905	33.003	-3.173	23.083	1.00 27.13
	ATOM	1890		LYS	905	28.039	-7.273	19.546	1.00 15.58
35		1891	Ö	LYS	905	27.251	-7.817	20.297	1.00 15.43
35	ATOM	1892	N	ILE	906	27.647	-6.620	18.466	1.00 15.71
	ATOM		CA	ILE	906	26.239	-6.554	18.086	1.00 15.74
	ATOM	1893		ILE	906	25.991	-5.423	17.030	1.00 14.76
	ATOM	1894	CB	ILE	906	24.527	-5.427	16.565	1.00 13.47
40	ATOM	1895	CG2		906	26.358	-4.051	17.611	1.00 13.40
40	MOTA	1896		ILE		26.021	-2.876	16.686	1.00 13.18
	ATOM	1897	CD1		906	25.800	-7.899	17.478	1.00 17.05
	ATOM	1898	C	ILE	906		-8.471	17.834	1.00 16.35
	ATOM	1899	0	ILE	906	24.759	-8.385	16.539	1.00 17.95
4.5	MOTA	1900	N	LEU	907	26.609		15.827	1.00 17.53
45	MOTA	1901	CA	LEU	907	26.348	-9.631		1.00 17.04
	MOTA	1902	CB	LEU	907	27.331	-9.787	14.659	1.00 13.32
	ATOM	1903	CG	LEU	907	27.338	-8.653	13.632	
	ATOM	1904	CD1		907	28.382	-8.885	12.557	1.00 12.28
	ATOM	1905	CD2		907	25.947	-8.531	13.029	1.00 13.60
50	ATOM	1906	C	LEU	907		-10.858	16.747	1.00 18.93
	ATOM	1907	0	LEU	907		-11.860	16.437	1.00 20.36
	ATOM	1908	N	SER	908		-10.805	17.868	1.00 19.02
	ATOM	1909	CA	SER	908		-11.947	18.772	1.00 19.60
	ATOM	1910	CB	SER	908		-12.099	19.407	1.00 19.21
55	ATOM	1911	OG	SER	908		-10.944	20.135	1.00 19.20
	ATOM	1912	Ċ	SER	908	26.027	-11.844	19.867	1.00 20.23
	ATOM	1913	ŏ	SER	908		-12.709	20.752	1.00 20.86
	ATOM	1914	N	GLY	909		-10.791	19.812	1.00 19.30
	ATOM	1915	CA	GLY	909		-10.603	20.817	1.00 18.22
60	ATOM	1916	C	GLY	909	24.533	-9.859	22.102	1.00 16.28
UU		1917	0	GLY	909	23.711	-9.712	22.987	1.00 16.09
	ATOM				910	25.773	-9.422	22.236	1.00 16.37
	ATOM	1918	N	LYS	310		2.746		

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	ATOM	1919	CA	LYS	910	26.166	-8.670	23.411	1.00	17.37
	ATOM	1920		LYS	910	27.665	-8.464	23.403	1.00	
	MOTA	1921		LYS	910	28.418	-9.703	23.684	1.00	
	ATOM	1922		LYS	910	29.860	-9.370	23.896	1.00	17.48
5	ATOM	1923		LYS	910		-10.534	24.482		17.90
5		1924		LYS	910	32.055		24.502	1.00	19.93
	ATOM	1925		LYS	910	25.472	-7.296	23.532	1.00	18.68
	ATOM ATOM	1926		LYS	910	25.250	-6.797	24.640	1.00	19.51
	ATOM	1927	-	VAL	911	25.219	-6.641	22.397	1.00	18.99
10	ATOM	1928		VAL	911	24.545	-5.341	22.396	1.00	17.49
10	ATOM	1929	-	VAL	911	25.501	-4.130	22.041	1.00	17.52
		1930	CG1		911	26.928	-4.550	22.019	1.00	15.48
	ATOM	1931		VAL	911	25.094	-3.412	20.788	1.00	
	ATOM	1931		VAL	911	23.379	-5.475	21.458	1.00	
45	ATOM	1933	_	VAL	911	23.504	-6.015	20.358	1.00	17.40
15	ATOM	1933		LYS	912	22.219	-5.032	21.896	1.00	17.96
	ATOM	1935		LYS	912	21.057	-5.210	21.072	1.00	19.32
	ATOM			LYS	912	20.189	-6.325	21.672	1.00	
	ATOM	1936 1937		LYS	912	19.261	-5.889	22.811	1.00	
20	ATOM			LYS	912	19.998	-5.297	24.030	1.00	
20	ATOM	1938 1939		LYS	912	19.509	-3.871	24.370		26.56
	ATOM		NZ	LYS	912	18.028	-3.782	24.457		27.08
	ATOM	1940		LYS	912	20.262	-3.943	20.903		19.72
	MOTA	1941	0	LYS	912	20.437	-2.985	21.651		19.46
25	MOTA	1942	N	PRO	913	19.463	-3.877	19.841	_	20.13
25	ATOM	1943	CD	PRO	913	19.437	-4.660	18.599		20.38
	ATOM	1944	CA	PRO	913	18.693	-2.665	19.683		20.09
	ATOM	1945	CB	PRO	913	18.174	-2.780	18.259	1.00	20.97
	ATOM	1946 1947	CG	PRO	913	18.127	-4.240	18.017		21.02
20	ATOM			PRO	913	17.555	-2.665	20.658		20.77
30	ATOM	1948	С	PRO	913	17.108	-3.719	21.120		20.82
	ATOM	1949	O N	ILE	914	17.100	-1.460	20.972		20.62
	ATOM	1950	CA	ILE	914	15.965	-1.262	21.846		18.90
	ATOM	1951 1952	CB	ILE	914	16.119	0.012	22.659		17.34
35	ATOM	1953	CG2	ILE	914	14.953	0.149	23.589		15.42
33	ATOM	1954		ILE	914	17.445	-0.022	23.418		16.40
	ATOM	1955		ILE	914	17.794	1.261	24.098		15.82
	ATOM	1956	CDI	ILE	914	14.823	-1.093	20.858		19.73
	ATOM	1957	Ö	ILE	914	14.946	-0.313	19.909		20.71
40	ATOM ATOM	1958	N	TYR	915	13.774	-1.908	20.995		19.80
40	ATOM	1959	CA	TYR	915	12.622	-1.823	20.105		19.03
		1960	СВ	TYR	915	12.194	-3.193	19.566		18.88
	ATOM ATOM	1961	CG	TYR	915	13.072	-3.773	18.505		18.76
	ATOM	1962	CD1		915	14.096	-4.640	18.832		19.63
45	ATOM	1963	CE1	TYR	915	14.923	-5.170	17.853		21.76
45	ATOM	1964		TYR	915	12.881	-3.457	17.173		19.64
		1965		TYR	915	13.698	-3.989	16.177	1.00	20.92
	ATOM ATOM	1966	CZ	TYR	915	14.721	-4.839	16.531		21.98
	ATOM	1967	OH	TYR	915	15.592	-5.314	15.577		25.00
50	ATOM	1968	C.	TYR	915	11.468	-1.273	20.882		18.68
50	ATOM	1969	Ö	TYR	915	11.340	-1.494	22.080		18.72
	ATOM	1970	N	PHE	916	10.621	-0.543	20.194		18.68
		1971	CA	PHE	916	9.456	-0.019	20.836		19.66
	ATOM	1972	CB	PHE	916	8.898	1.145	20.042		17.07
55	MOTA		CG	PHE	916	9.567	2.411	20.335		14.89
Ųΰ	ATOM	1973		PHE	916	9.377	3.034	21.561		16.16
	ATOM	1974		PHE	916	10.393	2.992	19.407		16.16
	ATOM	1975		PHE	916	10.010	4.225	21.854		14.78
	ATOM	1976			916	11.028	4.183	19.689		16.01
60	ATOM	1977	CE2	PHE	916	10.836	4.800	20.916		15.28
60	ATOM	1978 1979	C	PHE PHE	916	8.451	-1.148	20.868		21.60
	ATOM	1979	0	PHE	916	7.862	-1.434	21.910		22.04
	MOTA	1300	•	LUC	210	7.002	1.704			

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	N TOM	1981	N	HIS	917	8.300	-1.804	19.718	1.00 22.86
	ATOM	1982		HIS	917	7.354	-2.899	19.543	1.00 24.45
	ATOM ATOM	1983		HIS	917	6.549	-2.696	18.258	1.00 23.60
	ATOM	1984		HIS	917	5.921	-1.347	18.153	1.00 21.90
5	ATOM	1985	CD2		917	6.440	-0.153	17.787	1.00 21.97
5	ATOM	1986	ND1		917	4.614	-1.109	18.504	1.00 21.41
	ATOM	1987	CE1		917	4.350	0.178	18.360	1.00 22.05
	ATOM	1988	NE2		917	5.446	0.783	17.929	1.00 21.26
	ATOM	1989		HIS	917	8.077	-4.225	19.477	1.00 25.83
10	ATOM	1990	OT1	HIS	917	9.185	-4.257	18.908	1.00 27.53
	ATOM	1991	OT2		917	7.525	-5.225	19.988	1.00 29.26
	ATOM	1992	C1	DHT	920	27.685	5.199	4.565	1.00 13.59
	ATOM	1993	C2	DHT	920	26.814	6.485	4.636	1.00 12.55
	ATOM	1994	C3	DHT	920	25.484	6.280	3.944	1.00 12.58
15	ATOM	1995	03	DHT	920	24.904	7.249	3.448	1.00 11.99 1.00 13.18
	ATOM	1996	C4	DHT	920	24.887	4.964	3.857	
	ATOM	1997	C5	DHT	920	25.464	3.903	4.357	1.00 13.98 1.00 14.79
	MOTA	1998	C6	DHT	920	24.727	2.560	4.241	1.00 14.79
	ATOM	1999	C7	DHT	920	25.613	1.454	3.609	1.00 14.79
20	MOTA	2000	C8	DHT	920	26.955	1.303	4.359 4.279	1.00 13.34
	MOTA	2001	C9	DHT	920	27.708	2.656	4.279	1.00 14.56
	MOTA	2002	C10		920	26.943	3.876 2.525	4.830	1.00 14.73
	ATOM	2003	C11		920	29.161	1.344	4.192	1.00 14.11
	MOTA	2004	C12		920	29.951	-0.010	4.339	1.00 15.34
25	MOTA	2005	C13		920	29.194 27.784	0.212	3.680	1.00 15.67
	MOTA	2006	C14		920	27.178	-1.232	3.647	1.00 15.64
	MOTA	2007	C15		920	28.435	-2.118	3.310	1.00 15.37
	ATOM	2008	C16		920	29.679	-1.189	3.426	1.00 14.87
20	ATOM	2009	C17		920 920	29.107	-0.450	5.847	1.00 14.67
30	ATOM	2010	C18		920	26.781	3.770	6.524	1.00 13.94
	ATOM	2011	020		920	30.910	-1.918	3.981	1.00 16.20
	ATOM	2012	020	HOH	921	16.187	17.463	26.217	1.00 26.98
	ATOM	2013 2014	0	нон	922	19.878	17,183	14.290	1.00 13.49
35	ATOM ATOM	2015	ŏ	нон	923	18.473	14.908	14.407	1.00 6.52
33	ATOM	2015	ŏ	нон	924	29.144	18.703	11.673	1.00 37.40
	MOTA	2017	ŏ	нон	925	27.076	19.321	12.893	1.00 18.76
	ATOM	2018	ō	нон	926	23.789		9.649	1.00 33.78
	ATOM	2019	ō	нон	927	25.400	14.577	5.432	1.00 19.79
40	ATOM	2020	0	нон	928	23.015	12.473	12.245	1.00 14.03
. •	ATOM	2021	0	нон	929	25.209	14.445	2.442	1.00 19.95
	ATOM	2022	0	нон	930	34.235	16.490	0.235	1.00 41.09
	ATOM	2023	0	нон	931	31.687	16.720	1.143	1.00 22.88
	ATOM	2024	0	нон	932	26.451	12.094	2.237	1.00 8.25
45	MOTA	2025	0	нон	933	11.606	-0.191	-7.963	1.00 46.13
	ATOM	2026	0	нон	934			17.657	1.00 15.30
	MOTA	2027	0	нон	935	15.475	2.114	16.386	1.00 12.01 1.00 21.79
	MOTA	2028	0	НОН	936	8.514	-2.110	12.665 14.094	1.00 10.94
	MOTA	2029	0	НОН	937	23.094	0.783 -13.306	5.541	1.00 40.43
50	ATOM	2030	0	нон	938		-11.472	10.611	1.00 31.03
	ATOM	2031	0	HOH	939	26.933	-11.914	5.354	1.00 51.71
	MOTA	2032	0	нон	940	10.995	-6.843	16.294	1.00 29.91
	ATOM	2033	0	HOH	941	23.088	7.362	-10.811	1.00 30.10
55	ATOM	2034	0	HOH	942 943	26.671	9.139	-8.686	1.00 38.12
55	ATOM	2035	0	HOH	943	35.410	-8.438	-7.084	1.00 42.68
	ATOM	2036	0	нон нон	947	10.842	24.253	21.391	1.00 43.09
	ATOM	2037	0	НОН	947	15.704	21.095	27.707	1.00 54.35
	ATOM	2038 2039		НОН	949	1.671	16.382	5.866	1.00 24.50
60	ATOM ATOM	2039		НОН	950	8.009		8.572	1.00 36.16
00	ATOM	2040		НОН	951	29.490	17.190	30.961	1.00 56.26
	ATOM	2041		нон	952		-12.134	25.596	1.00 39.41
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	л том	2043	0	нон	953	42.457	5.523	7.132	1.00 28.93
	ATOM ATOM	2043	ŏ	нон	954	41.318	2.323	2.406	1.00 38.22
	ATOM	2045	Ö	нон	955	25.857	7.152	30.722	1.00 18.97
	ATOM -	2045	ŏ	нон	956	18.191	16.505	27.701	1.00 29.01
5	ATOM	2047	ŏ	нон	957	14.018	2.408	20.246	1.00 18.75
3	ATOM	2048	Ö	нон	958	14.651	4.006	17.873	1.00 21.70
	ATOM	2049	ŏ	нон	959	5.786	11.770	25.499	1.00 35.58
	MOTA	2050	ŏ	нон	960	2.694	19.497	9.834	1.00 25.35
	ATOM	2051	ō	нон	961	0.334	6.151	20.624	1.00 27.66
10	ATOM	2052	ŏ	нон	962	-2.677	2.639	17.420	1.00 35.67
10	ATOM	2053	ŏ	нон	963	0.868	8.543	25.138	1.00 43.49
	ATOM	2054	ŏ	нон	964	-8.085	7.667	23.358	1.00 40.82
	ATOM	2055	ŏ	нон	965	6.749	1.200	9.766	1.00 24.57
	ATOM	2056	ŏ	нон	966	-0.636	8.734	6.585	1.00 40.09
15	ATOM	2057	.o	нон	967	22.487	-4.734	14.335	1.00 28.04
13	ATOM	2058	ŏ	нон	968	18.615	17.070	7.167	1.00 23.83
	ATOM	2059	ŏ	нон	969	10.049	19.612	2.716	1.00 28.02
	ATOM	2060	ō	нон	970	26.829	21.030	22.736	1.00 25.40
	ATOM	2061	ŏ	нон	971	23.684	9.361	5.898	1.00 24.06
20	ATOM	2062	ō	нон	972	23.124	15.837	0.189	1.00 29.07
20	ATOM	2063	ō	нон	973	34.079	8.287	19.446	1.00 34.35
	ATOM	2064	Ō	нон	974	37.522	2.898	1.092	1.00 22.39
	ATOM	2065	Ó	нон	975	21.838	14.392	5.445	1.00 20.42
	ATOM	2066	ō	НОН	976	16.106	-10.859	0.784	1.00 48.09
25	ATOM	2067	0	нон	977	11.295	27.231	20.742	1.00 24.50
	ATOM	2068	0	нон	978	21.562	-7.923	18.100	1.00 34.94
	ATOM	2069	0	нон	979	41.647	-2.962	5.907	1.00 41.33
	ATOM	2070	0	нон	981	12.897	22.682	24.938	1.00 44.10
	ATOM	2071	0	нон	982	33.709	13.619	-5.931	1.00 26.84
30	ATOM	2072	0	нон	983	0.019	-4.834	14.164	1.00 36.91
	ATOM	2073	0	нон	984	39.563	3.365	-2.334	1.00 36.56
	ATOM	2074	0	нон	985	16.244	18.091	7.952	1.00 25.52
	ATOM	2075	0	нон	986	13.038	13.790	19.688	1.00 21.93
	ATOM	2076	0	нон	987	22.095	3.621	21.834	1.00 19.27
35	ATOM	2077	0	нон	988	2.516	2.235	3.905	1.00 30.91
	MOTA	2078	0	HOH	989	2.950	1.064	1.716	1.00 31.16
	ATOM	2079	0	нон	990	5.186	-1.082	5.207	1.00 26.66 1.00 28.42
	MOTA	2080	0	HOH	991	-0.310	15.229	24.529	
	ATOM	2081	0	нон	992	-6.181	9.210	18.935	1.00 37.84
40	ATOM	2082	0	нон	993	17.508	26.662	14.814 13.007	1.00 30.32 1.00 30.57
	ATOM	2083	0	нон	994	17.401	31.211	10.009	1.00 33.92
	ATOM	2084	0	нон	995	21.268	22.961 12.379	6.567	1.00 36.58
	ATOM	2085	0	нон	996	26.335 33.730	15.077	4.345	1.00 24.42
45	ATOM	2086	0	нон	997	28.576		-15.305	1.00 30.54
45	MOTA	2087	0	нон	998 999	33.926	-5 402	-13.979	1.00 48.01
	ATOM	2088	0	НОН		31.878		-10.283	1.00 38.45
	MOTA	2089	0	нон нон	1000 1021	30.673		18.256	1.00 34.16
	ATOM	2090 2091	0	нон	1022	35.035		15.084	1.00 37.70
50	ATOM ATOM	2092	ő	нон	1023	32.791		19.423	1.00 35.34
50	ATOM	2093	ŏ	нон	1024	22.587	-14.097	7.907	1.00 30.71
	ATOM	2094	ŏ	нон	1025	29.778		-0.255	1.00 24.68
	ATOM	2095	õ	нон	1026	25.904		24.176	1.00 16.80
	ATOM	2096	ő	нон	1027	33.066		7.455	1.00 20.84
55	ATOM	2097	ŏ	нон	1028	31.787		28.988	1.00 32.80
55	ATOM	2098	ŏ	нон	1029	27.029		13.994	1.00 20.01
	ATOM	2099	ő	нон	1030	20.499		16.384	1.00 31.66
	ATOM	2100	Ö	нон	1031	10.991	16.858	-1.085	1.00 30.58
	ATOM	2101	ŏ	нон	1032	7.904	10.344	-5.081	1.00 41.55
60	ATOM	2102	ŏ	нон	1033	12.570	3.398	-10.099	1.00 26.95
-	ATOM	2103	Ö	нон	1034	17.128		-10.962	1.00 22.24
	ATOM	2104	ŏ	нон	1035	17.056		-4.553	1.00 26.98

- 72 -6.595 1.00 25.24 11.020 0.892 нон 1036 2105 0 MOTA 1.00 27.44 -0.025 1037 3.092 -1.135 2106 0 HOH MOTA 5.653 1.00 34.54 24.006 35.765 HOH 1038 2107 0 MOTA 27.680 1.00 26.47 29.738 нон 1039 2108 0 MOTA 1.00 36.26 8.706 22.315 нон 1040 1.507 ATOM 2109 0 1.00 27.77 9.776 10.755 -4.751 2110 0 HOH 1041 MOTA 1.00 25.10 1.00 26.08 14.440 20.223 -3.560 нон 1042 2111 0 MOTA 30.147 -9.103 2.467 HOH 1043 2112 0 MOTA 1.00 28.96 28.518 -12.565 39.044 7.751 -5.152 нон 1044 2113 0 MOTA 17.961 1.00 38.02 1045 HOH 10 2114 0 ATOM 20.994 1.00 37.73 1046 37.030 10.428 нон 2115 0 MOTA 15.270 1.00 24.79 7.847 -2.227 0 HOH 1047 2116 MOTA 21.522 1.00 40.62 22.567 1.00 30.96 9.958 -5.351 1048 HOH 2117 0 MOTA 6.839 -6.928 нон 1049 2118 0

MOTA

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We claim:

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- 1. A crystal of an AR-LBD comprising:
 - a) an AR-LBD and an AR-LBD ligand or
 - b) an AR-LBD without an AR-LBD ligand;
- wherein said crystal diffracts to at least 3 angstrom resolution and has a crystal stability within 5% of its unit cell dimensions.
 - 2. The crystal of claim 1 wherein said AR-LBD has at least 200 amino acids.
 - 3. The crystal of claim 1, wherein said AR-LBD is the AR amino acid sequence 672 to 917 of rat AR
 - 4. The crystal of claim 1, wherein said AR-LBD is the AR amino acid sequence 672 to 917 of human AR.
 - 5. The crystal of claim 1 wherein the crystal comprises an AR-LBD and an AR-LBD ligand and the AR-LBD ligand is an agonist or antagonist, a partial agonist or partial antagonist, or a SARMs of the
- antagonist, a partial agonist or partial antagonist, or a SARMS of the AR-LBD.
 - 6. The crystal of claim 5 wherein the agonist is dihydrotestosterone.
 - 7. The crystal of claim 1 having all of the coordinates listed in Table A.
 - 8. The crystal of claim 1 wherein said crystal comprises mammalian AR-LBD protein.
 - 9. The crystal of claim 1 wherein said crystal comprises rat AR-LBD protein.
- The crystal of claim 1 wherein said AR-LBD ligand has the following unit cell dimensions in angstroms: a = 56.03 ± 5%, b
 = 66.27 ± 5%, c = 70.38 ± 5% and an orthorhombic space group P212121.
- A molecule or molecular complex comprising all or any part of the ligand binding site defined by structure coordinates of AR-LBD amino acids V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877 and F878 according to Table A, or a mutant or homologue of said molecule or molecular complex.

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- 12. The molecule or molecular complex of claim 11 wherein said mutant or homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said AR-LBD amino acids of not more than 1.5 Angstroms or 30% sequence identity with said AR-LBD amino acids.
- 13. A molecule or molecular complex comprising all or any part of the ligand binding site defined by structure coordinates of AR-LBD amino acids N705, Q711, R752, F764 and T877 according to Table A, or a mutant or homologue of said molecule or molecular complex.
- 10 14. The molecule or molecular complex of claim 13 wherein said mutant or homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said AR-LBD amino acids of not more than 1.5 Angstroms or 30% sequence identity with said AR-LBD amino acids.
- 15. A machine-readable data storage medium comprising a data storage material encoded with machine readable data, wherein the data is defined by the structure coordinates of an AR-LBD/AR-LBD ligand or ligand complex according to Table A or a homologue of said complex, wherein said homologue comprises backbone atoms that have a root mean square deviation from the backbone atoms of the complex of not more than 3.0Å
 - 16. The machine-readable data storage medium according to claim 15, wherein said AR-LBD/AR-LBD ligand or ligand complex is homologue having a root mean square deviation from the backbone atoms of said amino acids of not more than 2.0 Å.
- 17. A machine-readable data storage medium comprising a data storage material encoded with a first set of machine readable data comprising a Fourier transform of at least a portion of the structural coordinates for an AR-LBD/AR-LBD ligand according to Table A; which, when combined with a second set of machine readable data comprising an X-ray diffraction pattern of a molecule or molecular complex of unknown structure, using a machine programmed with instructions for using said first set of data and said second set of data, can determine at least a portion of the structure coordinates corresponding to the second set of machine readable data, said first set of data and said second set of data.

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- A binding site in AR-LBD for an AR modulator in which a portion of said ligand is in van der Walls contact or hydrogen bonding contact with any portion or all of residues V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898,
 I899, S900, V901, Q902, V903, P904 or I906 of AR-LBD according to Table A.
- The binding site according to claim 18 wherein the AR-LBD is a homologue or mutant with 25%-95% identity to residues V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906 of AR-LBD according to Table A.
 - 20. A method of obtaining structural information about a molecule or a molecular complex of unknown structure by using the structure coordinates set forth in Table A, comprising the steps of:
 - a. generating X-ray diffraction data from said crystallized molecule or molecular complex;
 - b. applying at least a portion of the structure coordinates set forth in Table A to said X-ray diffraction pattern to generate a three-dimensional electron density map of at least a portion of the molecule or molecular complex; and
 - c. using all or a portion of the structure coordinates set forth in Table A to generate homology models of AR-LBD or any other nuclear hormone receptor ligand binding domain.
 - 21. A computational method of designing an androgen receptor synthetic ligand comprising:
 - a. using a three dimensional model of a crystallized protein comprising an AR-LBD/AR-LBD ligand complex to determine

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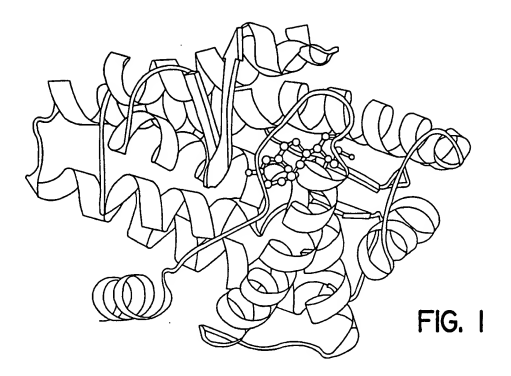
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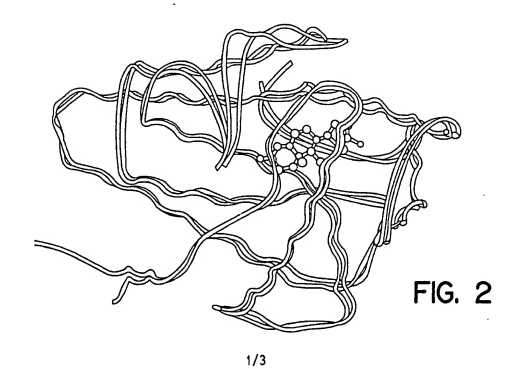
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- at least one interacting amino acid of the AR-LBD that interacts with at least one first chemical moiety of the AR-LBD ligand; and
- b. selecting at least one chemical modification of said first chemical moiety to produce a second chemical moiety with a structure that either decreases or increases an interaction between said interacting amino acid and said second chemical moiety compared to said interaction between said interacting amino acid and said first chemical moiety.
- 10 22. A method for identifying a compound that modulates androgen receptor activity, the method comprising any combination of steps of:
 - a. modeling test compounds that fit spatially into the AR-LBD as defined by structure coordinates according to Table A, or using a three-dimensional structural model of AR-LBD, mutant AR-LBD or AR-LBD homologue or portion thereof;
 - using said structure coordinates or ligand binding site as set forth in claim 18 to identify structural and chemical features;
 - employing identified structural or chemical features to design or select compounds as potential AR modulators;
- d. employing the three-dimensional structural model or the ligand binding site to design or select compounds as potential AR modulators;
 - e. synthesizing the potential AR modulators;
 - f. screening the potential AR modulators in an assay characterized by binding of a test compound to the AR-LBD;
 and
 - g. modifying or replacing one or more amino acids from AR-LBD selected from the group consisting of V685, L700, L701, S702, S703, L704, N705, E706, L707, G708, E709, Q711, A735, I737, Q738, Y739, S740, W741, M742, G743, L744, M745, V746, F747, A748, M749, G750, R752, Y763, F764, A765, L768, F770, M780, M787, I869, L873, H874, F876, T877, F878, L880, L881, V889, F891, P892, E893, M894, M895, A896, E897, I898, I899, S900, V901, Q902, V903, P904 or I906 of AR-LBD according to Table A.

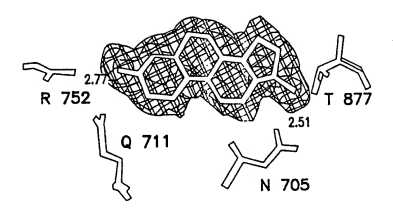
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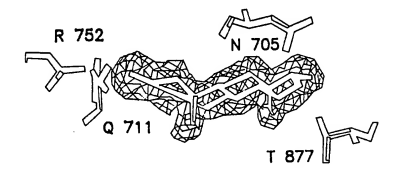
- 23. The method according to claim 22 wherein the potential AR modulator is from a library of compounds.
- 24. The method according to claim 22 wherein the potential AR modulator is selected from a database.
- 5 25. The method according to claim 22 wherein the potential AR modulator is designed de novo.
 - 26. The method according to claim 22 wherein the potential AR modulator is designed from a known agonist, partial agonist, antagonist, partial antagonist or SARMs.
- 10 27. The method according to claim 22 wherein the potential AR modulator is an agonist or partial agonist and AR activity is measured by translocation or unwinding or helix 12.
 - 28. The method according to claim 22 wherein the potential AR modulator is an antagonist or partial antagonist and AR activity is measured by translocation or unwinding or helix 12.
 - 29. An AR modulator identified by the method of claim 22.
 - 30. A method for treating prostate cancer comprising administering an effective amount of an AR modulator identified by the method of claim 22.
- 20 31. A method for treating an age related disease comprising administering an effective amount of an AR modulator identified by the method of claim 22.
 - 32. The method of claim 31 wherein said age related disease is osteoporosis, muscle wasting or loss of libido.





SUBSTITUTE SHEET (RULE 26)





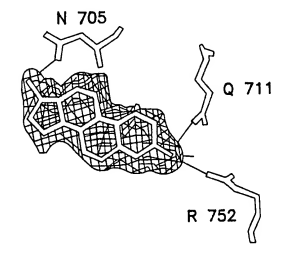


FIG. 3

SUBSTITUTE SHEET (RULE 26)

FIG. 4

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/28495

A. CLASSIFICATI N OF SUBJECT MATTER IPC(7) : GOIN 33/53										
US CL. :4	135/7.9	national classification and IPC								
According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED										
	ocumentation searched (classification system followed	by classification symbols)								
U.S. : 436/4,7.1,7.2,69.1; 436/501,63; 514/2										
Documentati searched	ion searched other than minimum documentation to	the extent that such documents are it	ncluded in the fields							
	ata base consulted during the international search (na BASE, MEDLINE, WPI, BIOTECH ABS, covering s									
c. poc	UMENTS CONSIDERED TO BE RELEVANT									
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.							
Y	US 5,411,981 A (GAILLARD-KELL) especially column 9, line 43, through c	30-32								
Y	US 5,434,176 A (CLAUSSNER et al.) column 8, line 49, through column 9,	30-32								
A	US 5,298,429 A (EVANS et al.) 2 document.	1-32								
Y	US 5,693,646 A (JONES et al.) 03 SUMMARY OF THE INVENTION SE	11-14, 18, 19, & 29								
Y	US 5,854,202 A (DEDHAR) 29 I document.	December 1998, see entire	11-14, 18, 19, & 29							
X Furt	her documents are listed in the continuation of Box (C. See patent family annex.								
Bpecial eategories of cited decuments: "I later document published after the international filing date or priority date and not in conflict with the application but cited to understand										
to be of particular subvance "X" decomment of particular subvance; the claimed investion cannot be conditioned nevel or cannot be conditioned nevel or cannot be conditioned to involve an investive step.										
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obvious to a person skilled in the art										
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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US00/28495

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
7	NAKADA et al. The Androgen Receptor Status of Neuroendocrine Cells in Human Benign and Malignant Prostate Tissue. Cancer Research. 01 May 1993, Volume 53, pages 1967-1970, see the entire document.	11-14, 18, 19, & 29
C	EKENA et al. Determinants of Ligand Specificity of Estrogen Receptor- α : Estrogen versus Androgen Discrimination. Journal of Biological Chemistry. 09 January 1998, Volume 273, Number 2, pages 693-699, see especially the abstract.	11-14, 18, 19. & 29
K 	MIYAMOTO et al. Promotion of agonist activity of antiandrogens by the androgen receptor coactivator, ARA70, in human prostate	11-14, 18, 19, & 29
Y	cancer DU154 cells. Proceedings of the National Academy of Sciences, USA. June 1998, Volume 95, pages 7379-7384, see the entire document.	30
X	SAI et al. An Exonic Point Mutation of the Androgen Receptor Gene in a Family with Complete Androgen Insensitivity. American Journal of Human Genetics. 1990, Volume 46, pages 1095-1100, see especially the abstract and the Introduction on pages 1095-1096.	11-14, 18, 19, & 29
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